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# Hydrogen applications and research activities in its production routes through catalytic hydrocarbon conversion

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**Abstract:** The statistical information on the share of hydrogen sector-wise consumption indicates that 95% of the total consumption is utilized in ammonia synthesis, petroleum refining processes and methanol production. We discuss how hydrogen is used in these processes and in several smaller-scale manufacturing industries. We also present the trend of hydrogen used as fuel, and as an energy carrier in fuel cells for generating electricity, powering hydrogen vehicles, as well as in aerospace applications. Natural gas caters for approximately half of the total hydrogen production resources. Therefore, the scope is emphasized on relatively recent developments in research activities related to the conventional catalytic hydrocarbon processing technologies for the production of hydrogen derived from natural gas (methane), which are steam methane reforming, partial oxidation of methane and autothermal reforming. Hydrocarbon decomposition is included due to its potential to be industrialized in the future, and its benefits of producing clean hydrogen without emissions of greenhouse gases and generating carbon nanofibers or nanotubes as by-products that have the potential in various emerging applications. Attention is given to the efforts toward achieving hydrocarbon conversion improvements, energy savings through thermally efficient operation and reduced operational costs through minimization or elimination of coke formation in the catalytic processes.

**Keywords:** autothermal reforming; hydrogen applications; hydrogen production; methane decomposition; partial oxidation; steam methane reforming.

## 1 Introduction

Hydrogen has been used mainly in petroleum refining processes such as desulfurization and cracking of oil, as an essential reactant, as well as in chemical industries as a raw material or syngas for synthesis of key chemicals; among others include methanol, dimethyl ether and ammonia (Chen et al. 2010). In addition, CO from the syngas (mixture of hydrogen and CO) is further chlorinated to produce phosgene, which is used as a starting point in the manufacturing other end-user chemicals including plastics, pesticides, dyes and herbicides (Chilcott 2007).

Based on historical data (Figure 1), it was indicated that just less than 50% of the total consumption of hydrogen in the USA in 1974 was in petroleum refining, followed by slightly above 35% in ammonia synthesis and up to 10% in methanol synthesis. This means that close to 95% of the total hydrogen consumption was made up by industrial-scale applications. The remaining pie was grouped together in miscellaneous uses (Dahiya and Chand 1987).

In 1975, a projection of hydrogen utilization for the year 2000 was made, where the industrial-scale applications would no longer be the leading consumers. Hydrogen consumption in the synthesis of chemicals (ammonia, methanol, etc.) and petroleum refining processes would only account for slightly above 40% of the total projected utilization. This is even less than half of the actual industrial-scale utilization in 1974, even though the total consumption was estimated to increase 22-fold from  $1.00 \times 10^5$  Btu (the unit was given as hydrogen energy in the statistical data).

The remaining hydrogen consumption was projected to be taken up by hydrogen used as synthetic fuels and energy, as well as for transportation and other energy-related uses. It was anticipated that hydrogen would be heavily utilized as a fuel in various industrial, commercial and domestic applications, as well as in the storage for use in automobiles and electrical power generation (Dahiya and Chand 1987).

Interestingly, after more than three decades, the global share of hydrogen consumption data in the 2000s (Ashik et al. 2015) showed that the utilization has still

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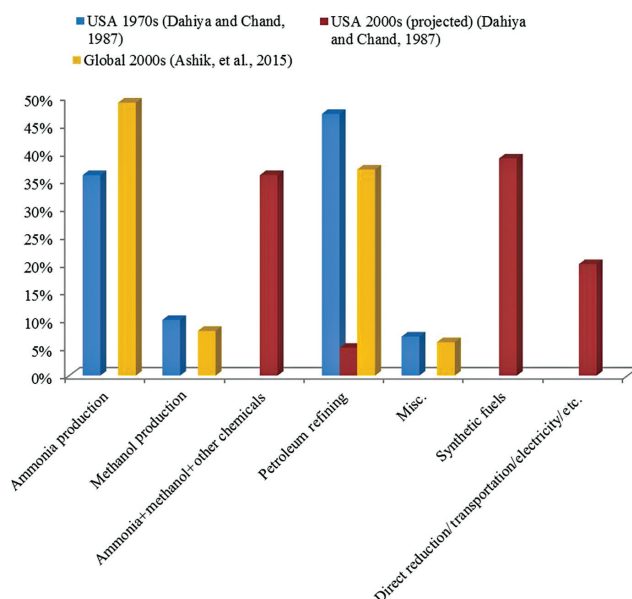


Figure 1: Share of hydrogen sector-wise consumption.

been dominated by industrial-scale applications. As presented in Figure 1, in comparison with the US hydrogen consumption data (1974), the only difference here is a swap of positions between ammonia production and petroleum refinery processes, but a similar trend is seen on hydrogen utilization in methanol production and miscellaneous uses.

Similarly in China, more than 95% of the total hydrogen consumption in the 2000s was utilized by the same top three large-scale sectors (i.e. synthesis of ammonia and methanol, and petroleum refining). Around 75% of the hydrogen was consumed for ammonia synthesis, at an ammonia annual production growth rate of 9%. Methanol output was also grown tremendously on yearly basis at the rate of as high as 39%, which increased accordingly the consumption of hydrogen in China. The increasing trend in the annual growth rate was also seen in fuel oil consumption at 7%, which subsequently led to a higher import volume of crude oil of high sulfur content, and hence resulted in an increase in the amount of hydrogen utilized in the petroleum refinery processes for the purpose of hydrofining and hydrocracking. Between 2000 and 2007, the total consumption of hydrogen in China increased at an annual growth rate of above 9%. The consumption increased by around 5 million tonnes in 2007 from 7 million tonnes in 2000 (Hewu et al. 2013).

The constituents of hydrogen utilization in the category of miscellaneous uses were not identified in the statistics. Based on the general applications of hydrogen presented in various literature, the miscellaneous

uses could include other industries such as electronics and semiconductors (Hewu et al. 2013), plastics, steel, ferrous and nonferrous metallurgy, food and edible oils processing (Dahiya and Chand 1987, Hewu et al. 2013) through a catalytic hydrogenation process, float glass and fine organic synthesis. Hydrogen is also used as part of the ingredients in the production of household cleaners (Navarro et al. 2007) as well as in the applications of aeronautics, astronautics and renewable-energy alternative vehicles (Hewu et al. 2013). Additionally, hydrogen is also utilized as an alternative fuel for energy and power in the industrial, commercial and domestic applications.

These miscellaneous uses of hydrogen do not greatly affect the total hydrogen consumption level (Dahiya and Chand 1987, Hewu et al. 2013), as their portion is relatively very small in comparison to the utilization in the top three large-scale industries. Taking hydrogen application as a transport fuel as an example, between July 2008 and August 2009, the passenger-carrying buses powered by hydrogen fuel cells in Beijing after traveling a total of 53,800 km consumed a total of only 5.33 tonnes of hydrogen (Hewu et al. 2013).

Not only hydrogen finds diversified applications, it is also versatile in its production technologies as it can be produced from various resources: fossil- and non-fossil-based. Approximately 48 million metric tonnes of the annual global industrial hydrogen production is based on fossil fuels (oil, natural gas and coal), which accounts for 95% of the total production, based on the statistics shown by the US Department of Energy from 1988 through to the 2000s (Figure 2) (Gaudernack, 1998, Ewan and Allen 2005, Navarro et al. 2007). Based on the consumption trend seen earlier, it is very interesting to note that the total hydrogen utilized by the large-scale industrial sectors is also around 95%. This means that by total demand-supply average,

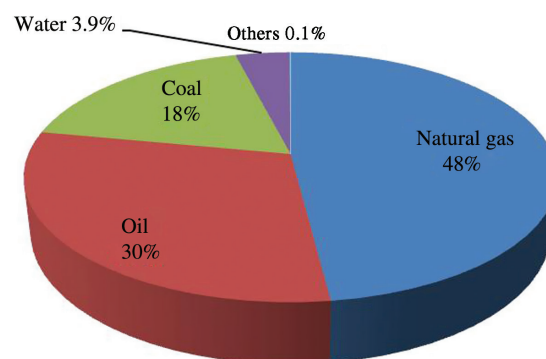


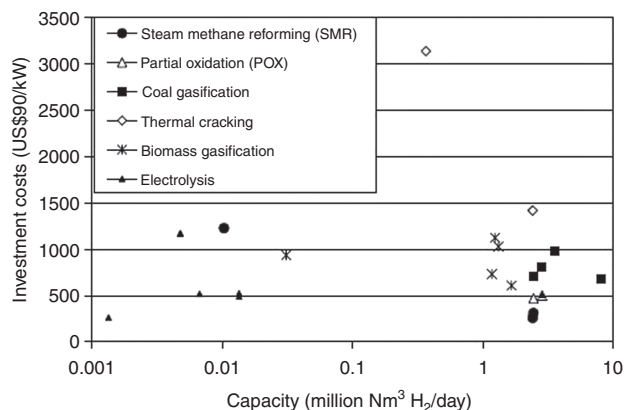
Figure 2: Sources distribution in total industrial hydrogen production of 500 billion N m³/year (Gaudernack 1998, Ewan and Allen 2005).

the requirement for hydrogen volume in these sectors and applications, in principal, can be fulfilled by the hydrogen produced from the fossil-based resources.

In hydrogen production, fossil resources can come from natural gas, oil and coal, whereas non-fossil resources can be derived from water, biomass and solid wastes. The infrastructures for the supply of these feedstocks can therefore be built according to what are available in the regions (Barreto et al. 2003). Different technologies have been developed to process different feedstocks in order to “extract” the hydrogen out. In general, natural gas is processed through the conventional catalytic operations. Light oil such as naphtha can be catalytically processed similarly as natural gas but some technologies are non-catalytic. Coal is processed through gasification (Conte et al. 2001).

Only a small proportion of hydrogen is produced from non-fossil resources. Biomass and solid wastes are processed through gasification similar to coal. Hydrogen derived from water, on the other hand, is produced through electrolysis, which can be powered by conventional fossil fuels, or alternative renewable power sources such as solar, wind, hydropower and nuclear power (Conte et al. 2001).

It is predicted that in the near future, fossil fuels will still be the primary resources for hydrogen production (Gaudernack 1998, Navarro et al. 2007). A survey conducted by Barreto et al. (2002) based on the published literature and public databases (Figure 3) shows that for the capacity of hydrogen production of beyond 1.0 million  $\text{Nm}^3/\text{day}$ , steam reforming of natural gas (methane) offers the least investment cost as compared to other technologies. Water electrolysis, whilst investment cost can also be low, is only capable of producing hydrogen in significantly small quantities.



**Figure 3:** Survey conducted by Barreto et al. (2002) on investment cost of hydrogen plants of various production capacities.

The hydrogen production technologies from natural gas in general have been around for decades and are highly efficient in large-scale operation (Hewu et al. 2013). Based on the above statistical data and investment costs analysis, the scope of this paper is therefore emphasized on the relatively recent development in the research activities related to the conventional catalytic hydrocarbon processing technologies for the production of hydrogen derived from natural gas (methane), which caters for close to half of the total hydrogen production resources. Before that, an overview of various applications of hydrogen will be presented.

## 2 Hydrogen applications

Hydrogen is traded on the market in liquefied form or compressed gas in cylinders, and some supplies are transferred via pipeline. Most hydrogen in chemical production and refinery processes is produced in and consumed by the same plants. A very large proportion of the world's hydrogen volumes are being used in the top three large-scale industrial processes (i.e. synthesis of ammonia and methanol, and petroleum refining). The analysis of the magnitude of the hydrogen volume consumed in these sectors will be briefly discussed. The hydrogen used in several smaller-scale manufacturing industries will also be presented.

We will also discuss the trend of hydrogen used as fuels for industrial, commercial and domestic consumptions. An overview of hydrogen used as an energy carrier in fuel cells for generating electricity and powering the hydrogen vehicles, as well as in spacecraft application, will also be presented to appreciate the use of this highly useful element that will shape our future.

### 2.1 Hydroprocessing in petroleum refineries

Hydrogen plays a role as a reactant in the refinery process where it is used in hydroprocessing. Hydroprocessing includes hydrocracking of heavy fractions, and removal of sulfur and other contaminants through hydrofining. The amount of hydrogen consumed by different refineries varies depending on the crude quality (i.e. level of sulfur and other contaminants contents) and characteristics (e.g. heavier crude requires a higher volume of hydrogen for hydrocracking) (Chernyi and Feigin 1968). Around 378,326,000 tonnes of crude was processed in China in 2007, consuming up to 1,700,000 tonnes of hydrogen, out

**Table 1:** Typical hydrogen consumption in various hydroprocessing stages in a typical refinery.

Hydroprocessing	H <sub>2</sub> consumption in wt% relative to crude oil feed
Hydrofining	
Straight run naphtha	0.01
Cracked naphtha	0.05–0.1
From low sulfur diesel feed to < 500 ppm diesel product	0.04
From high sulfur diesel feed to > 500 ppm diesel product	0.05
Hydrocracking	
Vacuum gasoil	0.5–0.8

Source: Castaneda et al. 2011.

of which 0.02 tonne was used in hydrocracking and 0.01 tonne in hydrofining, in every tonne of crude oil processed (Hewu et al. 2013).

Castaneda et al. (2011) reviewed approaches and techniques used to calculate hydrogen consumption required by a refinery, with the interest to optimize its utilization in both the hydroprocessing processes; the hydrofining and hydrocracking. The data presented in Table 1 are reproduced from Castaneda et al. (2011) to indicate typical hydrogen consumption in various processes in a typical refinery. Note that the large hydrogen consumption in hydrocracking of vacuum distillate increases the H/C ratio of the hydrocracker products relative to the hydrocracker feed, on top of utilization in hydrodesulfurization.

### 2.1.1 Hydrocracking

In the crude oil components, the higher boiling point fractions where the hydrocarbons are heavier are cracked into smaller chains upon heating. These smaller chains are hydrogen deficient upon cracking, and hence, they are instable. Hydrogen is added through catalytic hydrogenation (Dahiya and Chand 1987). The whole process is called hydrocracking, which is a simultaneous process of cracking and hydrogenation of hydrocarbons to break carbon-carbon bonds, and hence reduce molecular weight for higher yield production of refined and higher H/C ratio fuel products (Ramachandran and Menon 1998). The resulted fuel products have identical properties to those of gasoline (typical chain of C<sub>5</sub>–C<sub>10</sub>), which has the H/C ratio between 2.0 and 2.2 (Dahiya and Chand 1987).

In the event where the light crude supply drops, refiners have no other choices but to process more viscous and heavier crude oils that are of lower H/C ratios, which in

turn increases the requirement for hydrogen in the hydrocracking. The hydrogen requirement for hydrocracking varies depending on the feed composition, typically in the range of 1000 to 2500 standard cubic feet (SCF) per barrel of the crude processed (Dahiya and Chand 1987).

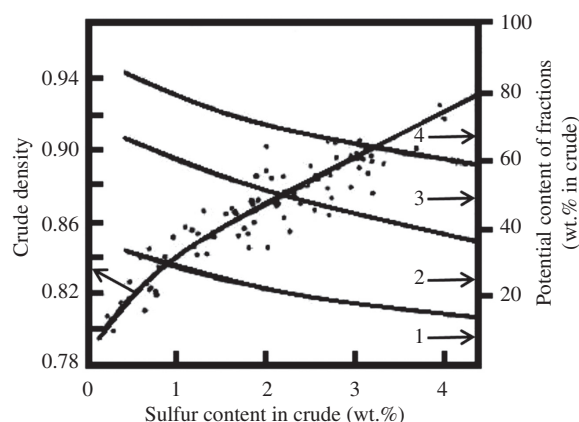
### 2.1.2 Hydrofining

Hydrofining is the technology used in refineries to upgrade oils with high impurities (Martínez and Ancheyta 2014). Sulfur, which is omnipresent in crude oil, is one of the main sources of particulates or soot emissions (Đukanović et al. 2013), as well as sulfur dioxide, a gaseous pollutant commonly associated with acid rain (Paez et al. 2015). In the hydrofining process, sulfur is removed from petroleum products in the form of hydrogen sulfide gas, which is then reacted with a base such as lime or lime stone to neutralize and stabilize the acidic compound to calcium sulfide. In general, at least 500 SCF of hydrogen is required for sulfur removal per one barrel of crude, depending on the sulfur contents (Dahiya and Chand 1987).

The current total sulfur content limit in transportation fuels such as gasoline and diesel in the EU, USA and some other parts of the world varies, where it is legislated at 10, 15 or 50 wt. ppm. Some countries still permit sulfur concentration limits of as high as 1000 ppm. Whilst compression ignition diesel engines are efficient in terms of fuel consumption, they adversely contribute to environmentally harmful emissions. Therefore, this has demanded the refineries to maximize the production yield of ultra-low sulfur diesel fuel (Đukanović et al. 2013). As for the gasoline, a review was made by Paez et al. (2015) based on TIER 3 of Transportation Impact Environmental Review of the US Environmental Protection Agency. TIER 3 establishes new standards in emissions cutting from light-duty vehicles, expecting to reduce sulfur content of gasoline from 80–30 ppm currently to 10 ppm.

In general, the sulfur content in a crude and the crude's density have a close relationship, and the density has a close connection with the crude's fractional compositions, as can be seen in Figure 4, where the data collected from Ural-Volga crudes were taken as an example. The light component consists of gas (fraction 1) and high boiling fraction (fraction 2) undergoes straight-run distillation to recover the products, called straight-run gasoline and diesel. The sulfur contained in the crude is distributed unevenly, where 60–70% sulfur concentrates in fractions 1 and 2. The distribution of sulfur in the secondary-processed products, called catalytically-cracked (fraction 3) and thermal-contact-cracked (fraction 4) gasoline and

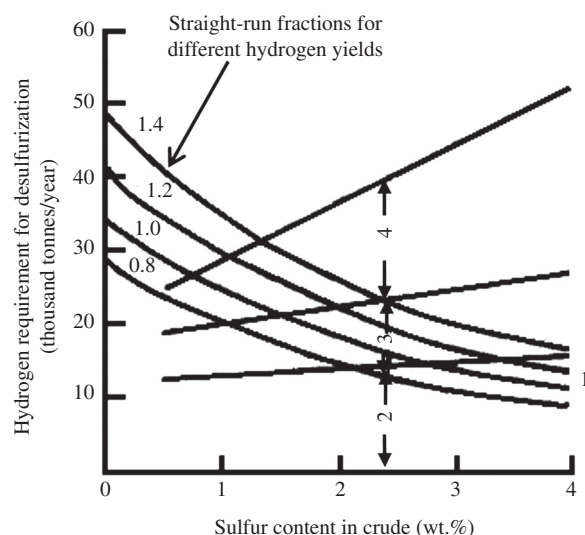




**Figure 4:** Left axis: sulfur content in crude as a function of crude density; right axis: crude fractions as a function of the crude density and sulfur content in the crude: (1) fraction with initial boiling point (IBP) of 180°C, (2) fraction with IBP 180–350°C, (3) fraction with IBP 350–500°C and (4) fraction with IBP 500°C (Chernyi and Feigin 1968).

diesel fuels, depends on the processing type (Chernyi and Feigin 1968).

Higher sulfur content in the crude gives a higher content of gasoline and diesel fuels obtained from both the straight-run distillation and the secondary processes, and hence, the desulfurization of these fuels requires a greater hydrogen volume. Figure 5 indicates the amount of hydrogen required in the hydrofining of the gasoline and diesel fuel, in relation to the sulfur content in the crude,



**Figure 5:** (1) Hydrogen production via catalytic reforming of straight-run fractions. (2, 3 and 4) Hydrogen consumption in hydrofining of gasoline and diesel fuels from straight-run fractions, catalytic cracking and thermal-contact cracking, respectively, in relation to crude's sulfur content (Chernyi and Feigin 1968).

for a refinery processing 12 million tonnes of crude. Note that hydrogen is also being produced via catalytic reforming of the straight-run fractions, shown by trend line 1. The trend in Figure 4 shows that the volume fraction of light components (straight-run fractions) in the crude decreases with the increase in the crude's sulfur content. With the decrease in the straight-run fractions, the hydrogen production via the catalytic reforming drops with the increase in the crude's sulfur content. On the other hand, the hydrogen consumption increases due to increased requirements in the hydrofining processes (Chernyi and Feigin 1968).

## 2.2 Ammonia synthesis

The Haber-Bosch ammonia production process started from a discovery of the chemical engineering foundations to synthesize ammonia from its elements in the laboratory by Fritz Haber, followed by an innovation by Carl Bosch who brought the production up to an industrial scale. Hydrogen is used in synthesis of ammonia ( $\text{NH}_3$ ) as raw material through an exothermic process carried out in a 3 to 1 molar ratio reaction with nitrogen. The reaction proceeds with high yields in the presence of iron catalysts.

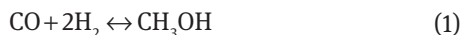
An interesting article featured in *Nature Geoscience* (2008) entitled "How a century of ammonia synthesis changed the world" (Erisman et al. 2008) discusses the significance of the Haber-Bosch process discovery in saving billions of people from hunger, due to the fact that fertilizers made from ammonia are mostly responsible in the expansion of agricultural activities for feeding the growing population in the world (Dahiya and Chand 1987).

Theoretically, the production of a tonne of ammonia requires an amount of approximately 0.18 tonnes of hydrogen (Hewu et al. 2013), which equals more than 30,000 SCF of natural gas (Dahiya and Chand 1987). In China, for example, an amount of 27,000,000 tonnes of ammonia was produced in 1997, which consumed 5,000,000 tonnes of hydrogen for synthesis. After more than 10 years due to the growing population, the numbers almost doubled up in 2008, where an amount of 51,589,000 tonnes of ammonia was produced using 9,411,900 tonnes of hydrogen (Hewu et al. 2013).

## 2.3 Methanol synthesis

Methanol is an important chemical used as a chemical storage carrier (Kobl et al. 2016, Xu et al. 2016), as well as in the transesterification process in biodiesel manufacturing

(Novinson 2004). Methanol synthesis from CO follows the reaction



Approximately 0.125 tonnes of hydrogen is required to synthesize 1 tonne of methanol. Over 20% of the world's methanol supply is met by China. In 1995, an estimation of 200,000 tonnes of hydrogen was consumed in producing 1,500,000 tonnes of methanol. Amazingly, in less than 15 years, the output of methanol in China amounted to 10,367,700 tonnes in 2008, consuming 1,310,000 tonnes of hydrogen (Hewu et al. 2013).

## 2.4 Other industries

Other than the three main industries mentioned above, hydrogen is also consumed in other industries as a reactant in various manufacturing and production plants, in small amounts. In addition, syngas can also supplement the demand for gasoline and fuels by conversion through Fischer-Tropsch synthesis (FTS), an emerging gas-to-liquid technology in the production of synthetic fuels.

### 2.4.1 Manufacturing and production industries

In the edible oils industry, hydrogen is used for hydrogenation of the edible oils to decrease the degree of unsaturated fats and increase the melting point such that the oils become a semi-solid at ambient temperature, for making margarine as an example. Hydrogenation, which takes place in the presence of nickel catalysts, also enhances the resistance to oxidation so that the food produced can be preserved longer (Dahiya and Chand 1987, Ramachandran and Menon 1998).

Whilst carbon is a good reducing agent to reduce metal oxides by combining with oxygen and forming gaseous oxides that escape away without further separation, hydrogen is an alternative reducing agent in metallurgical industry where carbon is undesirable. In tungsten manufacturing, for example, the presence of carbon makes the metal brittle, and hence, hydrogen is a better option (Dahiya and Chand 1987).

In electronics industry, wafer and circuit manufacturers use hydrogen as a reducing agent in the epitaxial growth of polysilicon, where silicon tetrachloride is reduced to silicon. In float glass manufacturing, the glass typically floats on a tin bath, and in order to prevent the oxidation of the molten tin bath, hydrogen is used as an oxygen scavenger in a mixture of 4% hydrogen in nitrogen (Ramachandran and Menon 1998).

### 2.4.2 Fischer-Tropsch synthesis

The premium pricing of gasoline and diesel fuels depends on their rating of research octane number (RON) and cetane number, respectively. Gasoline of a high RON would consist of highly branched alkanes, while diesel of a high cetane rating would consist of essentially linear alkanes. For example, *i*-octane (2,2,4-methyl pentane) has an RON of 100 and *n*-hexadecane has a cetane number of 100 (Dry 2002).

Engine fuels produced from crude oil distillation typically contain high amount of aromatics, organic sulfur and nitrogen. These compounds are environmentally undesirable. Furthermore, the aromatics decrease the cetane number, although at the same time increase the gasoline RON. Therefore, the diesel cut requires rigorous treatment with hydrogen to convert the aromatics to naphthenes as well as for hydrofining to bring down the sulfur and nitrogen contents. Despite the rigorous hydroprocessing, the cetane number is still not high enough when compared with that of linear alkanes (Dry 2002).

FTS coupled with some downstream treatment stages produces high-quality engine fuels as a result of low content of aromatics and zero content of sulfur. Fischer-Tropsch is a catalytic process that takes place at 200–300°C and 10–60 bars over iron and cobalt catalysts to convert syngas (mixture of  $\text{H}_2$  and CO) to a mixture of complex multicomponent hydrocarbons, ranging from linear and branched hydrocarbons to oxygenated products, with linear paraffins and  $\alpha$ -olefins as the main products. The synthetic sulfur- and aromatics-free FTS products are emerging as a complete substitute to the conventional engine fuels produced by crude oil distillation. At the very least, they can be blended together in meeting new quality regulations that will be more stringent in the near future (Van Der Laan and Beenackers 1999).

## 2.5 Hydrogen as an alternative fuel

Hydrogen is produced mainly from fossil fuel resources such as natural gas, oil and coal. These fossil resources are also used as fuels for generating various forms of energy such as thermal and electricity, by combustion with oxygen. Hydrogen is considered a clean alternative fuel to the fossil fuels, as the combustion of hydrogen with oxygen produces water, unlike the oxidation of fossils that leads to greenhouse gas emissions (Dahiya and Chand 1987). Presently, the EU as the world's most significant importer of fossil fuels is globally leading in the transition to renewable energy (Apak et al. 2012).

For decades, hydrogen has also been considered as an important alternative fuel (Apak et al. 2012). Hydrogen energy is in a chemical form and, therefore, for its utilization as a fuel in domestic, commercial, and industrial applications, or in automobiles and for generating electrical power, the chemical energy needs to be converted into usable forms of thermal, mechanical or electrical. Similar to other fossil fuels, the first step involves the conversion of hydrogen's chemical energy to thermal energy in the form of heat, via combustion with oxygen (Dahiya and Chand 1987).

### 2.5.1 High-temperature non-catalytic combustion for electricity generation

In electrical power plants, steam is generated by thermal or nuclear power, and the heat energy is converted in a turbine to motion power, to run the electrical generator. The oxidation of hydrogen as a fuel takes place at high temperature in the furnaces and combustors of the power plants. The fuel source from coal in the thermal power plants using coal-fired boilers can be supplemented with hydrogen, by replacing the oil support burners with hydrogen burners.

Some advantages of hydrogen as a substitute or a supplementary to fossil fuels (e.g. coal) include the elimination of (1) the corrosion and erosion problems in the boilers encountered when using coal due to the presence of impurities, and (2) the requirement to transport the coal to the power plant sites as the hydrogen gas can be supplied via underground pipelines.

In some electrical power plants, the electrical power produced can be stored for a later consumption through an interesting storing mechanism by having electrolytic plants installed at the site. The electrolysis process for producing hydrogen in the electrolytic plants is powered by the electrical power generated during peak hours. The electrolytically produced hydrogen is stored in metal hydride, as liquid hydrogen or as compressed hydrogen gas (Dahiya and Chand 1987).

### 2.5.2 Low-temperature catalytic combustion for small household and domestic appliances

Natural gas or liquefied petroleum gas that is being used for domestic and small-scale commercial applications, which account for about 20–30% of the primary energy consumption, can be substituted with hydrogen in the form of “town gas,” transported via pipelines. The town gas has a composition of 50–80% hydrogen with the

remaining balance being primarily carbon monoxide. Some of these applications apply combustion at ambient temperature in the presence of a catalyst, as the working principle. This catalytic combustion of hydrogen at room temperature is possible due to its low ignition energy (0.02 mJ). The exothermic reaction releases energy that is transferred via conduction and convection to heat an object or fluid that is then used for application in fuel-consuming devices for heating, cooking and electricity.

For instance, a catalytic burner made of catalyst impregnated heating panels provides the heat source for a flameless, safe cooking device. In household appliances such as fans and vacuum cleaners, the hydrogen fuel cells are used to generate the electricity needed for running them. In the application of heating homes that use hydrogen fuel, the need to exhaust the combustion product through a chimney is eliminated since the hydrogen combustion with oxygen in the air produces only water. Hence, good thermal efficiency can be achieved at a reduced construction cost (Dahiya and Chand 1987).

## 2.6 Emerging applications of hydrogen fuel cells for electricity generation, and in vehicles and spacecraft

Hydrogen is fast becoming the energy carrier of preference for a number of applications (Barreto et al. 2003), and hence, the demand of hydrogen for running fuel cells is expected to grow in the next decade (Navarro et al. 2007). Fuel cells technology is well developed and has reached a level of maturity. The working principle in the fuel cells involves a conversion of chemical energy directly to electrical energy by combining the hydrogen fuel gas with oxygen from the air, without involving a combustion, producing only water and heat as the by-products (Dincer 2002).

Different types of fuel cells have been developed with different characteristics; among others include operating temperature, heat availability, power density, as well as tolerance to thermal cycling and fuel impurities, which make each of the technologies suitable for a particular application. Dincer (2002) presented the working principles of a list of fuel cells that are the most commonly used and well developed, along with their respective applications. The fuel cells include solid oxide electrolyte fuel cells, molten carbonate fuel cells, phosphoric acid fuel cells, proton exchange membrane fuel cells, unitized regenerative fuel cells, alkaline fuel cells (AFCs) and direct methanol fuel cells. The applications of fuel cells cover a wide range of sizes from hand-held computing and communication devices powered by miniature, circuit-like

and easily “rechargeable” fuel cells, to ultra-light, very efficient and low-polluting cars powered by inexpensive low-temperature fuel cells, and finally fuel cells used as small distributed electricity generators (Barreto et al. 2003).

The United States enacted energy legislation under the Energy Policy Act of 2005 (Holt and Glover 2006), which was passed due to concerns raised in security of energy and the quality of the environment. According to Lokey (2007), the section on hydrogen in Title VIII in the Act provides a direction toward hydrogen as a substitute to the declining petroleum global supplies. In the provisions, an authorization of \$3.3 billion for FY 2006–2010 was allocated for research and development activities on a hydrogen and fuel cell, especially with increased attention to hydrogen fuel and fuel cell vehicles following the Hydrogen Fuel Initiative announced at the State of the Union Address in January 2003 (Holt and Glover 2006).

### 2.6.1 Electricity generation

The increasing interest in hydrogen energy has seen hydrogen as an energy carrier being manufactured for electricity. As presented in Apak et al. (2012), Seoul, Korea, is an example of a capital city that takes seriously the use of hydrogen energy for electricity. The aim in their new climate change master plan looks at expanding the country’s renewable energy share from 1.5% to 20% between 2007 and 2030, which includes contribution of hydrogen energy of close to half of this share.

Fuel cells are regarded as clean power producers, as there are no emissions of the acid rain or smog producing pollutants, as they do not utilize the combustion of fossil fuels. In addition, they are considered as efficient power producers as they generate electricity in a simple step and at significantly lower temperatures relative to electricity production at conventional power plants, where the combustion of fossil fuels takes place at a high temperature for heat generation. A few steps are involved in conventional power production that prior to the generation of electricity, the produced heat energy is first converted to mechanical energy (Dincer 2002). In hydrogen-air fuel cells, the efficiency of the hydrogen chemical energy conversion to electrical energy can go as high as 80% (Dahiya and Chand 1987).

### 2.6.2 Vehicles and forklifts

Andrews and Shabani (2012) envisioned hydrogen’s role as a key in a sustainable energy economy for not only

longer-term storage in electricity grids, but also in the range of road and rail vehicles as well as marine ships and air transport. Due to its non-polluting properties, hydrogen is used in internal combustion engines as fuel, similar to engines operated with petrol and diesel (Apak et al. 2012). An analysis by Lin et al. (2013) revealed that hydrogen vehicles, including hydrogen internal combustion engines and hydrogen fuel cells, possess great potential to reduce reliance on petroleum and thereby reduce greenhouse gas emissions. The hydrogen-fuel-cell-powered electric vehicles have the potential to offer a sustainable solution to the renewable resources transportation market. The hydrogen system as a whole from distribution, refueling and on-board storage offers superiority over batteries recharged with grid power (Dincer 2002).

In transportation applications, hydrogen is supplied for consumption in fuel cells by several ways. One of the options is through the hydrogen production at central locations via small-scale natural gas steam reforming, biomass or coal pyrolysis or gasification, or water electrolysis powered by renewable solar, wind, hydro or geothermal energy, followed by its storage on board the vehicle as a gas or a cryogenic liquid, or stored in a solid carbon. Another example of an option is through the use of methanol as a hydrogen carrier where the methanol is chemically broken into hydrogen and carbon dioxide by a catalyst on the fuel-cell’s membrane (Dincer 2002).

The largest opportunity for fuel cell technology in the transportation market is in the light-duty automotive applications. Most of the players in the automotive industry have at least introduced fuel cell vehicle programs in their portfolios. The prototype of vehicles powered by hydrogen fuel cells have been developed and launched by many of them (Dincer 2002, Navarro et al. 2007). A list of existing commercialized hydrogen fuel cells for transportation by various automakers can be found in Conte et al. (2001) for further readership.

In the heavy-duty transportation sector, the attractiveness of the hydrogen fuel cell systems, especially in meeting a zero emissions target, was discussed in a study carried out by Ally et al. (2015) based on the trends of energy use and emissions in the road transport sector in Australia. More interestingly, the study also discussed the capability of the hydrogen fuel cell in meeting the requirements for range, duty cycle or payload higher than that of battery-only vehicles. Andrews and Shabani (2012) shared the same view where hydrogen fuel cell vehicles were identified as the transportation mode for medium or long distance trips, while plugin battery electric vehicles are reserved for short trips.



Operating the forklifts is an equally interesting application of fuel cells. In comparison with the battery-powered forklifts, the ones operated by fuel cells offer various advantages that include higher productivity due to elimination of battery changing that is time consuming. Additionally, they can be refueled very quickly at the hydrogen fueling stations around the site. Unlike the forklifts powered by batteries, fuel cells have a significant advantage of maintaining a constant voltage that does not drop toward the end of a shift or in cold sites. A typical fuel-cell forklift design has a fuel-cell stack that acts as the main power source. The power source is supplemented by the energy stored in a battery or super-capacitor during peak demand (Elgowainy et al. 2009).

### 2.6.3 Space

Hydrogen in liquid form is used in the aerospace industry. Cryogenic liquid hydrogen is used as a lightweight fuel to propel space shuttles, rockets and aircrafts. Additionally, since the 1960s, the electrical systems on the spacecraft have been powered by hydrogen fuel cells in which the process also produces water for the crew's drinking consumption (Dahiya and Chand 1987, Wu and Leu 2014). Cecere et al. (2014) presented the historical journey from the aerospace industry applications of hydrogen since its discovery, through to its current and future technology advancements in turbojet, ramjet, scramjet and rocket engines, for further readership.

The type of fuel cell used in the space missions by NASA is the AFC, which has a power generating efficiency as high as 70%. Although AFCs are among the oldest and simplest type of fuel cells, their commercial applications are very costly. An AFC is made of porous carbon plates deposited with a catalyst as the electrodes, with potassium hydroxide as the electrolyte. The hydrogen gas fuel combines with the hydroxide ions at the anode to produce water vapor, which results in left-over electrons. The produced water and the electrons form the hydroxide ions at the cathode, and these ions are recycled back to the anode for interaction with hydrogen, and the cycle repeats (Dincer 2002).

### 2.6.4 Issues pertaining to infrastructure requirements for enabling widespread hydrogen-powered appliances

Despite all the attempts to increase the popularity of hydrogen fuel cell vehicles, an interesting thought was

raised by Krumdieck and Dantas (2008). According to them, the process of substituting fossil fuels with renewable fuels is not simple and straightforward. Based on the feasible-sustainability concept, the whole network for the solution must take into account changes in land use, infrastructure provision and technology capability. Renewable energy transport, such as electric cars and hydrogen fuel cell cars whilst presently getting the hype as the solution to sustainable transport, requires challenging infrastructure considerations. The issues presented in this review are meant to put the widespread use of hydrogen-powered appliances into perspective without the intention to address the approaches and solutions on overcoming them.

In order to facilitate widespread adoption of hydrogen-powered appliances, the infrastructure to produce, transport, store and distribute the hydrogen must be well developed. Competitive small-scale decentralized hydrogen generation units need to be established, with distribution systems that are highly reliable and can meet the power quality standards need to be put in place (Barreto et al. 2003). An identification of the future locations of hydrogen plants will need to take technical, economic and political aspects into considerations due to the large space requirement. Land availability will depend on its topography and may compete with meeting other purposes such as vegetation for instance. In exploiting wind energy as another example, there are not many regions in the world that are suitable for large-area wind energy utilization. Most of the suitable regions are mountainous coastline, and hence, the installations of wind power system become impractical for both economic and ecological reasons (Nitsch and Voigt 1988).

## 3 Research activities in hydrogen production technologies

### 3.1 Water electrolysis

At a smaller scale, hydrogen can be produced by splitting water into oxygen and hydrogen via electrolysis, which accounts for approximately 3.9% of the total hydrogen production, as seen in Figure 2 (Gaudernack 1998, Ewan and Allen 2005). Increased attention on considering renewable sources has also opened up pathways for utilization of hydropower, and solar and wind energy for hydrogen production. Based on the report by National Research Council and National Academy of Engineering of the National Academies entitled "The Hydrogen

Economy: Opportunities, Costs, Barriers, and R&D Needs” (Board on Energy and Environmental Systems National Research Council 2004), the Department of Energy of the United States has listed the future direction on research and development focus in electrolysis through initiatives toward reducing cost and enhancing the efficiency of the system through integration between electrolyzers with power derived from wind and solar energy. Hydropower serves mainly to replace the fossil fuels in the electrical power plants to meet the huge industrial demand for electricity, and hence, only a small portion is utilized for electrolytic hydrogen production (Nitsch and Voigt 1988).

Wind energy collected in remote windy regions presents an economically unlimited energy source. Solar energy on the other hand requires a consideration on the factors such as the availability of large amounts of radiation and the condition of the identified geographical areas, so that the construction and the operation of the solar plants can be done at low costs in order for them to be economical. Not only the total amount of radiation is crucial, the shares of the diffuse and direct radiation must also be taken into account in the feasibility evaluation of building a solar plant (Nitsch and Voigt 1988).

Hudson et al. (2009) reviewed methods that employ solar energy, namely, photovoltaic (PV) and photoelectrochemical (PEC). The PV technique is a process utilizing PV electricity generation where the electricity is generated first, followed by the electrolysis. Whilst being feasible, its efficiencies are limited. PEC, on the other hand, is a single-step process where solar light illuminating semi-conducting photoelectrode produces electrons and holes, and hence dissociates the water. Wu and Leu (2014) presented several techniques of recovering hydrogen from water electrolysis in their review. Among others include high-temperature decomposition and the photo-biological process.

Marshall's group has been actively producing novel research in this area. One of their works demonstrated proton exchange membrane water electrolysis that offers various advantages in comparison to the traditional alkaline technologies in terms of energy efficiencies and hydrogen production rates (Marshall et al. 2007). Another study exhibited the use of electro-catalytic cobalt oxide layers to enhance the efficiency of water electrolysis cell (Mellsop et al. 2014).

### 3.2 Gasification

As seen in Figure 2, up to 18% of the total hydrogen production comes from coal, while a very small portion of 0.1%

production is categorized as others, which could include biomass- and solid waste-based resources. Both of these fossil and non-fossil resources are consumed to produce hydrogen through gasification (Gaudernack 1998, Ewan and Allen 2005, Navarro et al. 2007).

Gasification is the main pathway of producing hydrogen using coal. Other than gasification, coal coking is another means of producing hydrogen. In gasification, coal is burned in the presence of oxygen (air) or vapor and converted into gaseous products where hydrogen is the main component, at a high temperature and a constant pressure, or under an elevated pressure. In coking, coal is heated to produce coke and coke oven gas by-product, at temperatures of 900–1000°C without the presence of air. Typically, in every 1 tonne of coal, approximately 300–350 N m<sup>3</sup> of coke oven gas that has hydrogen content of up to 60% can be produced from the coking process. The remaining components of coke oven gas include CO, unsaturated hydrocarbons, CO<sub>2</sub>, O<sub>2</sub> and N<sub>2</sub> at varied concentrations. The coke oven gas has a heat value of 17–19 MJ/N m<sup>3</sup>, which makes it a good fuel candidate in high-temperature industrial furnaces or as a town gas. In China, the majority of the hydrogen produced from coal is used for the synthesis of ammonia and methanol. For example, of 7.13 million tonnes of hydrogen produced from coal in 2007, up to 78.5% was used for ammonia production, and slightly less than 15% was used for methanol production (Hewu et al. 2013).

Biomass has also received a lot of attention as the resource for hydrogen production. An overview and perspective of a system called biomass integrated gasification combined cycle with steam gasification was presented by Pang and Li (2006), for electricity and thermal energy generation from wood. Among other key areas for research work in this subject includes gasification performance optimization and capital cost minimization. For instance, a study undertaken by Pang's group (Saw et al. 2012) on steam gasification of mixtures of biomass sources demonstrated an increase in H<sub>2</sub> and CO production as well as improvement in the H<sub>2</sub>/CO ratio in the fluidized bed gasifier when wood pellet mass was added with increasing load of dried sewage sludge called biosolids.

Hosseini and Wahid (2016) presented an overview of the hydrogen production technologies using biomass resources via biological and thermochemical processes. The biological conversion process includes photo-fermentation and water biophotolysis using algae. The thermochemical process is found to be superior due to its lower cost and higher efficiency compared to the biological process. It was also found that supercritical water gasification (SCWG) is a more cost-effective process compared

to steam gasification. The SCWG process utilizes highly moisturized biomass directly in the gasification process, which eliminates a drying stage that is typically costly. In addition, the hydrogen is produced at high pressure, and, therefore, a relatively smaller energy amount is required to pressurize the gas in the storage tank.

### 3.3 Conventional catalytic hydrocarbon processing technologies

Based on the statistical data in Figure 2 and the investment costs analysis in Figure 3, the production of hydrogen derived from natural gas (methane) through conventional catalytic hydrocarbon processing technologies accounts for slightly below half of the total global hydrogen production, and hence, becomes the main focus of discussion in this review. In addition, these catalytic processes are also applicable in processing petroleum oil components such as naphtha as the resource, which accounts for 30% of the total production.

The most commonly used catalytic hydrocarbon conversion technologies for large-scale hydrogen production are the conventional steam methane reforming (SMR), partial oxidation of methane (POX) and autothermal reforming (ATR) (Gaudernack 1998, Holladay et al. 2009, Ciambelli et al. 2011). The average release of  $\text{CO}_2$  to the atmosphere of  $0.6 \text{ N m}^3$  in every  $\text{N m}^3$  of hydrogen produced from fossil fuels (natural gas, oil and coal) corresponds to a global  $\text{CO}_2$  release of approximately 500 Mt in a year, which is a considerably large amount of greenhouse gas.  $\text{CO}_2$  sequestration would reduce substantially the amount of the release but this comes with an exorbitant price. However, the cost increase is expected to be moderate in the case of hydrogen production from natural gas (Gaudernack 1998).

Hydrocarbon decomposition is included in this review not only because it has the potential to be industrialized in the future, but also because the technology produces clean hydrogen and eliminates the CO emissions. The objective of the methane (natural gas) decomposition process is to crack the hydrocarbon into hydrogen and carbon. The decomposition was originally employed for the production of carbon black used for energy storage, while hydrogen was produced as a by-product to supplement the fuel for the process (Gaudernack 1998, Muradov 2002).

Muradov (2002) presented the working principles of different types of hydrocarbon decomposition technologies currently being developed in a laboratory scale, among others include thermal (pyrolytic), plasma, photocatalytic and thermocatalytic decomposition. In this

review, we are interested in the last type of hydrocarbon decomposition technology due to its advantage in generating carbon nanofibers (CNFs) or nanotubes as by-products that have the potential in various emerging applications (Teo et al. 2003).

Even though the catalytic natural gas processing technologies for hydrogen production have been around for many decades, there is always room for improvement in the form of improving the hydrogen output, minimizing the total energy consumption and reducing operating cost by avoiding problems such as carbon deposition in the process.

#### 3.3.1 Steam methane reforming

This section presents an introduction to the commercial SMR process, and research efforts in understanding the catalytic process over various transitional metal catalysts, on different carrier materials, as well as process modifications to make the process effective. The thermodynamics and kinetics of the reactions involved, together with the transport processes of the process gas and molecules over the catalyst particles, are a separate topic of discussion, and hence not covered in this review.

Steam reforming is a principal process for hydrogen and synthesis gas production in the industry, where more than 80% of global ammonia production is based on the hydrogen produced by steam reforming (Mondal and Chandran 2014) prior to synthesis with nitrogen. The first step of the SMR process involves the reaction of methane with steam at  $697\text{--}827^\circ\text{C}$  to produce syngas (reaction (2)), which is a mixture of hydrogen and carbon monoxide. Water gas shift reaction (WGSR) then follows, where the CO produced in the first reaction reacts with steam and form hydrogen and carbon dioxide (reaction (3)). The overall SMR reaction is represented by the combined reaction (reaction (4)) (LeValley et al. 2015).

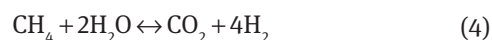
SMR:



WGSR:



Combined SMR and WGSR:



The research undertaken in SMR has been quite fundamental, where most work involved catalytic performance study to improve the reaction conversion.

Different interpretations of reaction kinetics presented in the literature were reviewed by Jones et al. (2008), and they appeared to follow a general consensus of the first-order reaction with respect to methane.

Nickel-based catalysts are not just traditionally used for steam reforming (Jones et al. 2008), but have also been the most commonly used due to their adequately high activity and considerably lower cost compared to other metal catalysts (Wu et al. 2013). A study was conducted on SMR kinetics at varying Ni%-Al<sub>2</sub>O<sub>3</sub> to investigate the effect of the Ni content (Kim et al. 2015). From the analysis, the activity was found to be higher at higher Ni loading, undoubtedly attributed to the higher number of Ni metal particles.

Nickel catalysts nevertheless possess the drawback of being prone to deactivation due to carbon deposition from hydrocarbon cracking, even when being operated at the conditions where steam-carbon ratios are anticipated to be thermodynamically outside the regime of carbon formation (Wu et al. 2013), hence, inviting a number of studies to address and identify the order of reactivity of a series of transition metals for superiority comparison. Ru, Rh, Pd, Ir and Pt are among the noble metals that have been demonstrated to provide good activity for SMR, but they are not justified cost-wise for use in industrial reformers (Jones et al. 2008).

Jones et al. (2008) conducted the experimental work to rank the steam reforming activity for Ni, Pt, Ru, Rh, Ir and Pd supported on ZrO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub>, at 773 K and atmospheric pressure, which is a condition away from equilibrium. The activities were correlated to the uniformity and homogeneity of metal catalysts dispersion, by measuring the adsorption energies of hydrogenated atoms adsorbed on the pure metal surfaces. The approach was considered vital due to the entropic effects that are significant at the high steam reforming reaction temperatures. The technique allowed thermodynamic parameterization of the kinetic model for determination of the activities of the pure metals in terms of turn-over frequencies, which is the parameter to quantify catalyst activity measured by  $\text{mol}_{\text{reactant\_reacted}}/\text{s.g}_{\text{catalyst}}$  (Fogler 1999). The relative catalyst reactivity was in agreement with other earlier publications in the order of Ru > Rh > Ni > Ir > Pd > Pt from higher to lower activity at similar dispersion (Jones et al. 2008).

It has also been observed that catalytic research in SMR has evolved and progressed from activity investigations on pure metal to metal alloy catalysts. The addition of a second metal to the “traditional” nickel catalyst has been demonstrated to significantly modify the properties of nickel particles as a result of bimetallic surface alloy formation. Wu et al. (2013) were inspired to compile the

studies on the use of bimetallic catalysts, by investigating catalytic activity and catalyst deactivation by carbon formation over modified nickel surfaces through formation of an alloy with a second metal from a series of noble and other metals including Au, Ag, Sn, Cu, Co, Mo, Fe, Gd and B. In general, Ni/second-metal bimetallic catalysts had been proven to exhibit greater catalytic performance compared to monometallic nickel catalysts in terms of activity, resistance to coke deposition and metal particles’ sintering and hydrogen selectivity.

Besides bimetallic alloy catalyst systems, studies have also been performed on composites of metal mixtures to improve SMR performance. For instance, a composite catalyst of nickel, cerium, zirconium and zinc was synthesized in a relatively recent study that integrated SMR with WGS in the effort to bring down production costs (LeValley et al. 2015). Interestingly, the composite catalyst was designed to provide activity for both the SMR and WGS. A significant finding was noted on the addition of zinc that resulted in reduced surface area, and hence, the activity of the SMR reaction, but on the other hand increased WGS activity, based on the observed improvement in carbon dioxide selectivity, hydrogen production and resistance to carbon deposition. Additionally, this composite catalyst system is potentially valuable in low-temperature SMR investigations, a condition which is less favorable for carbonaceous gases emission, based on the promising methane conversion and CO<sub>2</sub> selectivity performance at a reaction temperature of 650°C.

The fundamental catalytic study on SMR has focused on the effect of the metal catalyst addition as a promoter or surface modifier of the nickel catalyst with the aim of improving energy efficiency and reaction performance. Investigations have also included modifications to the catalyst support, which acts as the carrier, to meet the same objectives.

The carrier has its own specific role in the nickel catalyst activity. Ni on its own is not active, until it is distributed on a carrier. A carrier enhances the surface area of Ni and ensures that small Ni crystallites will not bind together and form larger crystals (thermal sintering). The carrier is designed to have a certain porosity to allow gas access to the entire available Ni surface. It is a stable, inert and mostly ceramic-based material where no activity for accelerating any undesired side reactions should be expected, and it must also not react chemically with any constituents of the process gas.

Having a high fusion point or high thermal stability is also an important criterion for selecting a carrier, as SMR is highly endothermic and takes place at very high temperatures (typical inlet temperature of 450–650°C



and outlet temperature of 700–950°C). However, carriers are typically acidic and the selection should also take into account their level of acidity. The acidity can cause hydrocarbon cracking activity that leads to coke (carbon) formation.

Antzara et al. (2016) evaluated SMR performance in chemical looping using a NiO-based catalyst supported on a series of oxygen transfer materials:  $\text{ZrO}_2$ ,  $\text{TiO}_2$ ,  $\text{SiO}_2$ ,  $\text{Al}_2\text{O}_3$  and  $\text{NiAl}_2\text{O}_4$  at a low temperature of 650°C and a steam-methane ratio of 2 : 1. It was found that NiO/ $\text{SiO}_2$  and NiO/ $\text{TiO}_2$  had low activity with less than 50% of the methane being converted in the beginning and eventually dropped to less than 10% methane conversion after 2 h on stream as they deactivated rapidly. NiO/ $\text{ZrO}_2$ , NiO/ $\text{Al}_2\text{O}_3$  and NiO/ $\text{NiAl}_2\text{O}_4$  exhibited good catalytic activity with an initial methane conversion of more than 80%, but only NiO supported on  $\text{ZrO}_2$  demonstrated good stability. On the other hand, NiO supported on  $\text{Al}_2\text{O}_3$  and  $\text{NiAl}_2\text{O}_4$  exhibited high deactivation that led to a drop in methane conversion to 59% and 63%, respectively, at the test end. This superior stability of  $\text{ZrO}_2$  is consistent with an earlier research work by the same group of researchers, investigating NiO supported on  $\text{Al}_2\text{O}_3$  and  $\text{ZrO}_2$  (Shen and Lua 2015), in experiments carried out in multiple cycles of reduction/reforming at 650°C and reoxidation at 850°C.

In the following study presented by Kho et al. (2016), which was carried out recently, a promising low-temperature SMR operation was demonstrated on titanium dioxide-supported and silica-supported nickel catalysts, Ni/ $\text{TiO}_2$  and Ni/ $\text{SiO}_2$ . The former offered stability and effective hydrogen production at 500°C, with methane conversion of 45% at a steam-methane ratio of 3 : 1. This is in alignment with the study by Antzara et al. (2016) that demonstrated a similar level of activity at 650°C, which indicates that an even lower operating temperature could be achieved in SMR using nickel catalyst supported on titanium dioxide without affecting its level of activity, by increasing the steam-methane ratio. The activity performance at low temperature was attributed to stronger support interaction with nickel species. In comparison to nickel supported on conventional inert oxide (silica), the reaction for hydrogen production was more active via WGSR, instead of SMR, due to a weaker nickel-support interaction.

One significant research area in SMR of late is the modification of the catalyst support substrate to a monolith. As commonly practiced, the SMR reaction takes place in metal tubes containing a packed bed of typically Ni/ $\text{Al}_2\text{O}_3$  particulate. The slow and highly endothermic reaction requires large amounts of heat supplied via external firing of reformer tube, in which

conventionally the heat transfer is ineffective due to poor convection from the reactant species to the active catalyst sites. Monolithic support substrates have been introduced where the catalysts are deposited on it and heat transfer efficiency is hence improved via the enhanced thermal conductivity of the monoliths, transferring the heat from the tube wall to the catalyst sites. This efficient process leads to process intensification where a higher reaction rate is achieved in a smaller reactor at a significantly lower pressure drop. Furthermore, the monolith has improved the mechanical integrity of the catalytic system. Giroux et al. (2005) and Heck et al. (2001) presented a discussion on the advantages of this promising catalytic support system.

Other than the fundamental catalytic performance investigations through modifications of nickel catalyst surfaces and catalyst supports, knowledge in chemical engineering has also been applied in modifying the so-called conventional SMR process design as well as in designing new reactor configurations, for reaction improvement.

Zhu et al. (2015) presented a novel work applying CaO sorption enhanced SMR (SESMR) for thermal promotion of hydrogen production. The system was combined with chemical looping combustion (CLC) for  $\text{CO}_2$  capture, instead of the commonly used furnace. The design was developed through Aspen Plus simulation. CLC employment served to function as the provider of the necessary heat for SESMR, while at the same time achieving inherent separation of  $\text{CO}_2$  without extra consumption of energy. The developed configuration exhibited advantages over the existing commercial SMR process, most notably on hydrogen purification, without the need for an additional process to remove impurities, while being more energy efficient. CO and  $\text{CO}_2$  concentrations were decreased alongside with the progress of the reaction, from 0.2% and 2.4% down to ppm level.

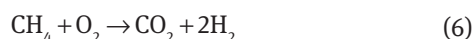
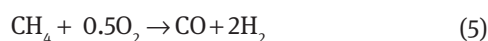
With the objectives to reduce energy consumption and carbon formation as well as to improve reaction performance, Kyriakou et al. (2016) investigated the feasibility of SMR at low temperatures (450–650°C) by employing a Ni-BZCY72/BZCY72/Cu proton conducting membrane reactor (MR). The MR had the flexibility to perform as a typical reformer operation by conducting the test under open-circuit conditions as the first step. Membrane-based operation took place in the same system, when hydrogen produced from the first step was electrochemically transported from the anode made of Ni-BZCY72 to the Cu cathode. The membrane operating mode was observed to have demonstrated further improvement in methane conversion and hydrogen yield, after the first open-circuit step.

Another novel reactor design was developed by Choi et al. (2016) by elimination of the use of catalysts, hence, eliminating the risk of deteriorating reaction performance due to catalyst deactivation. In addition, it was reported that hydrogen yield of higher than 70% was achieved at atmospheric pressure, in the developed microwave system, composed of mainly a 2.45 GHz microwave plasma torch and a plasma nozzle, that generated a steam plasma at which SMR took place. The need for catalysts was eliminated as the high-temperature plasma flame enhanced the reaction rate while the steam microwave plasma excited the activity of the reacting species.

SMR research has been focused on fundamental work in various types of metal catalysts aimed at achieving an energy-efficient operation at high reaction performance and minimal carbon deposit, which typically include the examination using various mono- and bi-metal catalysts, as well as modifications of the catalyst support and process engineering design. Table 2 summarizes the work reviewed in this paper.

### 3.3.2 Partial oxidation of methane

In the POX process, the fuel is combusted with oxygen to produce syngas, following exothermic reactions (5) and (6) (Muradov 2002, Dedov et al. 2015). POX is favorable in comparison to steam reforming, for the use in Fisher-Tropsch and methanol synthesis (Dedov et al. 2015):



In general, the partial oxidation process can take a wide range of hydrocarbon feedstocks. Any heavy hydrocarbons including heavy residual oils and coal can be processed by non-catalytic partial oxidation at high temperatures of around 1100–1500°C. On the other hand, the catalytic partial oxidation of light hydrocarbon fuels such as natural gas and naphtha takes place at significantly lower range temperatures of around 600–900°C (Muradov 2002).

The production of hydrogen by means of partial oxidation can be derived from various hydrocarbon molecules as well as biomaterials. Catalyst activity in the partial oxidation of bioethanol was studied by Kraveva et al. (2015) over xNiAlZn catalysts at various Ni loadings, while Wang et al. (2016) investigated partial oxidation of a mixture of phenol, acetic acid and naphthalene in super-critical water.

As far as POX is concerned, active investigations have been carried out by researchers on various transitional

metal catalysts to understand their performance and conditions toward achieving the main objectives of improving  $\text{H}_2$  and CO (syngas) selectivity, increasing hydrogen yield and eliminating carbon deposition. Efforts include the addition of support materials to the metal catalysts, such as mixed perovskite-like oxides to minimize carbon formation, the use of monolithic support to improve selectivity and reactor modifications to increase product yield.

In two very recent works investigating catalytic activity and coke formation at varied operating parameters, nickel catalysts deposited on nanocrystalline  $\text{CeO}_2$  and  $\text{Al}_2\text{O}_3$  supports were used in POX (Peymani et al. 2016) and propane (Peymani et al. 2017), respectively. The nickel catalysts prepared by an impregnation technique that exhibited a high surface area, and a mesoporous structure with very uniform size distribution, were varied in their loading content, where the highest catalytic activity was observed on 10% Ni/ $\text{CeO}_2$  and 7.5% Ni/ $\text{Al}_2\text{O}_3$ . With a rise in the reaction temperature, the amount of carbon accumulation increased slightly in the case of  $\text{CeO}_2$  in methane partial oxidation, but decreased slightly in the case of  $\text{Al}_2\text{O}_3$  on propane. Nonetheless, the deposited carbon amount decreased with the increase in the oxygen-methane molar ratio from 0.25 to 0.75 in both cases, together with an improved methane conversion in the case of methane partial oxidation. Other factors such as varied calcination and reduction temperatures during the catalyst pretreatment were also studied, to understand their effect on the hydrocarbon conversion. It was found that the methane conversion dropped with the increase in the calcination temperature, while propane conversion improved with the increase in the reduction temperature.

In the efforts of mitigating carbon deposition that can deactivate the catalysts, initiatives on studying the effect of a second metal have been made. For instance, very little carbon was formed during catalytic partial oxidation reaction tests using Ni-Ru bimetallic catalyst as compared to when single metal nickel catalysts were used (Velasco et al. 2015). Another example is that the addition of CoO to  $\text{ZrO}_2$  supported NiO catalyst, which has also proven to significantly reduce carbon deposition (Choudhary et al. 1998). Mixed perovskite-like oxides (Dedov et al. 2015, Palcheva et al. 2015) when being added to the catalysts served to improve catalyst activity due to their ability to generate well-dispersed metal catalyst particles during reduction, stabilizing the small metal catalyst clusters.

In reaction performance investigations, Diehm and Deutschmann (2014) and Figen and Baykara (2015) presented studies of catalytic methane partial oxidation using monolithic catalysts. Diehm and Deutschmann (2014) studied performance of the reaction over palladium- and

Table 2: Areas of study in steam methane reforming and their corresponding potential benefits.

Area of study	Investigation	Finding	Achievement	Investigator
Catalyst	Composite catalyst of nickel, cerium, zirconium and zinc	Development of composite catalyst (by co-precipitation of mixed metal nitrates), simultaneously active for both SMR and WGSR	Important step in emerging toward low-temperature SMR of 650°C	LeValley et al. 2015
Catalyst support	NiO-based catalyst on $ZrO_2$ , $TiO_2$ , $SiO_2$ , $Al_2O_3$ and $NiAl_2O_4$ at 650°C and $H_2O-CH_4$ ratio of 2 : 1	NiO/ $ZrO_2$ exhibited high catalytic activity (initial $CH_4$ conversion of > 80%) and good stability NiO/ $TiO_2$ exhibited low activity (< 50% $CH_4$ conversion) and rapid deactivation Ni/ $TiO_2$ exhibited good stability, with $CH_4$ conversion of 45% due to strong metal-support interaction at lower temperature Ni/ $SiO_2$ demonstrated lower activity due to weaker metal-support interaction	Low-temperature SMR operation at 650°C (energy saving)	Antzara et al. 2016
Catalyst support	Ni-based catalyst on $TiO_2$ and $SiO_2$ at 500°C and $H_2O-CH_4$ ratio of 3 : 1	Ni/ $TiO_2$ exhibited good stability, with $CH_4$ conversion of 45% due to strong metal-support interaction at lower temperature Ni/ $SiO_2$ demonstrated lower activity due to weaker metal-support interaction	Lower temperature of SMR operation at 500°C using Ni/ $TiO_2$ (more energy saving, improved catalyst stability than operation at 650°C)	Kho et al. 2016
Catalyst support	Monolithic support	Higher reaction rate in a smaller reactor at a lower pressure drop and improved mechanical integrity of the catalytic system	Process intensification	Heck et al. 2001, Giroux et al. 2005
Process design	Aspen Plus simulation of thermal promotion using CaO sorption enhanced steam methane reforming (SESMR) combined with chemical looping combustion (CLC) for $CO_2$ capture	CLC captured $CO_2$ while provided heat for SESMR without additional energy required $H_2$ with purity as high as 92.6% Improved overall energy efficiency to 85.5% from 68% (in the conventional SMR process)	Cost saving due to an energy-efficient system and removal of the additional hydrogen purification step	Zhu et al. 2015
New reactor configuration	Ni-BZCY72/BZCY72/Cu proton conducting membrane reactor	Highly active Ni-BZCY72 electrode at low $H_2O-CH_4$ ratio (< 2), with $CH_4$ conversion close to equilibrium (80% conversion) and 77% $H_2$ yield at 650°C	Low-temperature (450–650°C) SMR operation with high $CH_4$ conversion and $H_2$ yield	Kyriakou et al. 2016
New reactor configuration	Activity of reacting species excited by a 2.45 GHz steam microwave plasma, and hence, eliminated the use of catalyst	$H_2$ yield of higher than 70% achieved at atmospheric pressure	Eliminated catalyst use, hence coke deposition problem while maintaining good $H_2$ yield	Choi et al. 2016

rhodium-coated alumina monoliths, while Figen and Baykara (2015) investigated reaction performance over Co, Co-Ni, Co-Ru, Co-Ni-Ru and Ni catalysts supported on cordierite monoliths. Neagoe et al. (2016) demonstrated more effective oxidation of methane in  $O_2/Ar$  mixtures by observing higher selectivity of both CO and  $H_2$  at 900°C and 20 bar using Pt supported on iron-chromium alloy (FeCrAlloy) woven metal, when compared with commercial Pt-gauze or Pt/Rh gauze.

Through a new design on reactor configuration, a numerical simulation study by Chen et al. (2010) demonstrated an achievement in process intensification where the reaction was run over a rhodium catalyst bed embedded at the center of a Swiss-roll structure reactor. The simulation demonstrated improvement in methane conversion and hydrogen yield of higher than 10% in POX through preheating of reactants using waste heat recovered from hydrogen gas. This achievement was made possible by the exchanger design in the Swiss-roll structure where the number of turns in the reactor was increased and the gas hourly space velocity (GHSV) was brought down for the reactants to interact effectively with the catalyst bed.

The performance of methane partial oxidation in an MR has also attracted researchers to investigate the options of membrane material for the purpose of process intensification and hydrogen production improvement. Chibane et al. (2011) conducted theoretical analysis of Pd-MR performance through periodic feed composition and carrier gas, which was proven to have produced significantly superior level of methane conversion and hydrogen recovery, in comparison to steady state feed. Song et al. (2015) performed an experimental comparison between different membrane materials, namely, perovskite oxides membrane and conventional dense membrane. The former exhibited mixed ionic and electronic conductivities, indicating higher stability and better performance as it possesses high oxygen permeation flux, and hence, higher methane conversion efficiency. A model developed based on a multi-layer asymmetric oxygen-permeable membrane by Shelepova et al. (2016) was simulated to determine the process parameters such as temperature and gas flow rates, and membrane parameters such as pore diameter of porous layer and thickness of porous layer, at which methane conversion was the highest and syngas selectivity was the best.

Air Products and Chemicals in collaboration with the Department of Energy of the United States through Ceramtec have developed ceramic ion transport membranes for separating oxygen of high purity from the air and conducting the ionic oxygen at high temperatures, particularly economical for the use in partial oxidation processes due to high level of oxygen purity and possibility

of high flux transfer of oxygen. The technology exhibits a process intensification initiative where air separation and the reacting system are uniquely integrated, attractive for compact design that has the potential to reduce significantly the construction cost of large-scale units.

An article by Dyer et al. (2000) presented this membrane technology development and cited its corresponding registered patents for further reference. In the early 2000s when the concept was introduced and proven at a laboratory scale, some technical hurdles have yet to be overcome in the efforts to commercialize the units in an industrial scale. The challenges include the lack of design that can integrate successfully the engineering processes with the ITM, the unavailability of an economical technology for the ceramic membrane fabrication, and the undeveloped potentially stable ITM materials that can meet the required performance characteristics. Fast forward to 2014, Miller et al. (2014) presented the advancement of the original ITM technology for syngas production through the addition of porous layers on the membrane where it was demonstrated that the highest oxygen fluxes were achieved due to an increase in the membrane's surface area. Whilst the technology offers huge potential for cost reduction in syngas production processes, the technical hurdles have not been completely overcome in materializing the technology application at an industrial scale, and in fact the technology is no longer pursued by Air Products.

An interesting model was developed by Dashliborun et al. (2013) using a one-dimensional steady state non-isothermal model to investigate POX performance using a side feeding air injection reactor (SFR) and compare with a fixed bed reactor (FBR) and an MR. The simulated results demonstrated enhancement of the reaction performance with the increase in the number of air injections in the SFR. With only a single air injection, the outlet temperature in the SFR already showed a reduction of close to 200 K from 1269 K in the FBR. Hydrogen selectivity improved to 53% when the number of air injections in the SFR was increased up to 3, from 50% in the MR. This revealed that the performance was equivalent, and hence, the MR, which was associated with high process cost, could now be potentially substituted. The finding also led to a creation of controllable operating temperature and enhancement of hydrogen selectivity, which would potentially eliminate the problems of high operating temperature in FBR, hence, reduction in the operational cost through energy saving.

Another interesting reactor configuration in the experimental study of POX is the microchannel reactor (Makarshin et al. 2015). It was demonstrated that the temperature profile along the FeCrAlloy microchannel plate



length deposited with Ni-Pt/LaCeZrO catalyst was not flat. Exothermic combustion of methane into  $\text{H}_2\text{O}$  and  $\text{CO}_2$  took place at the front edge of the plate, while endothermic reactions of methane steam and dry reforming occurred on the remaining part of the plate, yielding CO and  $\text{H}_2$ . One microchannel plate demonstrated methane conversion of 82% and CO selectivity of 60% at a GHSV of  $820,000 \text{ h}^{-1}$  and under thermocycling of up to  $900^\circ\text{C}$ . It was concluded that hydrogen production is proportional to the number of microchannel plates.

Other than attempts in meeting the objective to improve hydrogen yield, there have also been other engineered solutions to study POX presented in the literature for energy efficiency. One of them is the modeling study of POX on a methane-steam blend in a flow reactor of a given length conducted by Starik et al. (2015). Hydrogen-rich syngas production was achieved through oxygen activation by electric discharge. This plasma-chemical method exhibited significant achievement in the ratio of energy returned to energy consumed.

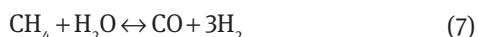
Wang et al. (2015) presented a new concept to increase the efficiency level of syngas production from solid oxide  $\text{H}_2\text{O}/\text{CO}_2$  co-electrolysis that previously had been found to require high energy input. The introduced concept combined co-electrolysis with POX to efficiently generate syngas on both sides of the electrolyzer, hence, demonstrating lower electric input and cathodic co-electrolysis polarization resistance, as compared to conventional co-electrolysis.

In summary, efforts have been made in the subject of methane partial oxidation to meet the objectives of minimizing carbon formation, increasing yield and selectivity, and improving energy efficiency. Summarized in Table 3 are the initiatives put forward in the work on POX reviewed in this paper.

### 3.3.3 ATR of methane

The complete ATR includes exothermic POX, the WGSR and the endothermic SMR reaction. The major chemical reactions are as follows (Lee et al. 2005):

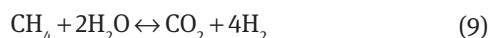
Methane steam reforming reaction:



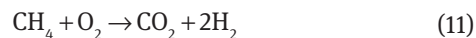
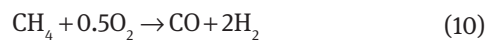
WGSR:



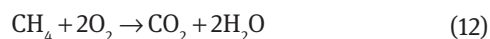
Coupling of steam reforming and WGSR:



Methane partial oxidation reaction:



Methane complete oxidation reaction:



Yan et al. (2013) presented the study on characteristics of methane ATR to generate hydrogen using a thermodynamic equilibrium constant method, developed by MATLAB programming. Their investigation results showed that reaction (7) is prone to reverse at low temperature. By manipulating the steam-methane and air-methane ratios, the inflection point temperature for the reaction to move forward was successfully identified.

Their findings give useful information on molar ratio regulation. For example, at a molar ratio of steam to methane fixed at 2, an increase in the air-methane molar ratio brought up the inflection point temperature. On the other hand, fixing the air-methane molar ratio at the value of 1 would maintain the inflection point temperature in the range of 700–800 K. At temperature exceeding 1000 K, an opposite result was observed where an increase in the air-methane molar ratio brought down the production level of hydrogen. The optimized parameters for easy occurrence of methane ATR were demonstrated at a temperature of 1000 K, an air-methane molar ratio of 1 and a steam-methane molar ratio of 2. The result also led to the finding that selectivity of  $\text{H}_2/\text{CO}$  can be manipulated by regulating the steam-methane molar ratio (Yan et al. 2013). The air-methane ratio is one of the parameters manipulated in the attempt to achieve better conversion in the secondary reformer unit in a typical ammonia plant (Al-dhfeery and Jassem 2012).

Further manipulation of variables to find the optimized condition was simulated by Yan et al. (2015) in ATR of methane in a microreactor, using CHEMKIN. The effects on methane conversion, carbon deposition and product yield were investigated by varying the temperature, oxygen-carbon ratio, steam-carbon ratio and catalyst activity density. The findings are useful for the development of a mobile fuel cell.

Nickel has been the active metal used in the catalytic investigations of methane ATR. A study on the effect of nickel content supported on a high specific surface area nanocrystalline  $\text{MgAl}_2\text{O}_4$ , on the catalytic activity was demonstrated by Mosayebi et al. (2012). The highest methane conversion was observed on 15% Ni/ $\text{MgAl}_2\text{O}_4$  compared to other lower nickel loading amounts in all the

**Table 3:** Modifications of existing systems to improve partial oxidation of methane process.

Area of study	Investigation	Finding	Achievement	Investigator
Catalyst loading amount optimization	Varied nickel loading supported on high surface area $\text{CeO}_2$ and alumina for $\text{CH}_4$ and $\text{C}_3\text{H}_8$ partial oxidation, respectively	Optimized catalytic activity observed on 10% Ni/ $\text{CeO}_2$ and 7.5% Ni/ $\text{Al}_2\text{O}_3$ at an $\text{O}_2$ - $\text{CH}_4$ molar ratio = 0.25–0.75	Deposited carbon decreased with an increase in the $\text{O}_2$ - $\text{CH}_4$ molar ratio	Peymani et al. 2016, 2017
Catalyst modification	Addition of Ru to Ni catalyst as a promoter at thermodynamically favourable coke formation condition (at low $\text{O}_2$ - $\text{CH}_4$ and $\text{H}_2$ - $\text{O}-\text{CH}_4$ ratios)	No appreciable amount of carbon on Ru; small amounts of carbon formed on bimetallic Ni-Ru compared to Ni catalysts at $\text{O}_2$ - $\text{CH}_4$ = 0.2, $\text{H}_2$ - $\text{O}-\text{CH}_4$ = 0.3, reaction temperature = 700°C	Carbon formation effectively overcome by the use of Ru-containing catalysts	Velasco et al. 2015
Catalyst modification	Addition of CoO to NiO catalyst at a ratio of 1, supported on $\text{ZrO}_2$ , $\text{ThO}_2$ and $\text{UO}_2$	Reduction in carbon deposition as compared to using NiO catalyst solely at $\text{O}_2$ - $\text{CH}_4$ = 0.51, reaction temperature = 700°C	Reduction in carbon deposition but a decrease in conversion and selectivity	Choudhary et al. 1998
Catalyst modification	Application of mixed perovskite-like oxide catalysts	$\text{CH}_4$ conversion ~ 90%, syngas selectivity > 90% with $\text{H}_2$ /CO ~ 2 due to finely dispersed $\text{Nd}_2\text{O}_3$ , CaO, cobalt oxides and cobalt metal from $\text{NdCaCo}_{0.96}$ decomposition at POX conditions of $\text{CH}_4$ - $\text{O}_2$ = 1.8–4.5 and 850–960°C	Improvement in overall reaction performance	Dedov et al. 2015
Catalyst modification	Application of mixed perovskite-like oxide catalysts	Rh perovskite catalyst with $x$ = 0.25 gallium exhibited the highest catalytic activity of 83% $\text{CH}_4$ conversion at 600°C (versus $x$ = 0.1 and 0.4), contributed by reducibility of $\text{La}_{0.75}\text{Sr}_{0.25}(\text{Fe}_{0.8}\text{Co}_{0.2})_{1-x}\text{Ga}_x\text{O}_{3-3.8}$ perovskite materials	Improvement in overall reaction performance	Palcheva et al. 2015
Catalyst support modification	Monolithic support	Staged $\text{Pd}/\text{Al}_2\text{O}_3 + \text{Rh}/\text{Al}_2\text{O}_3$ coated honeycomb monoliths versus single-sliced $\text{Pd}/\text{Al}_2\text{O}_3$ and $\text{Rh}/\text{Al}_2\text{O}_3$ coated honeycomb monoliths at an $\text{O}_2$ - $\text{CH}_4$ ratio = 1	General demonstration of comparison in reaction performance improvement	Diehm and Deutschmann 2014
Catalyst support modification	Monolithic support	No $\text{H}_2$ yield increase observed on staged catalyst compared to single-sliced catalysts of the same length		
Catalyst support modification	Monolithic support	Syngas formation on Rh followed direct route, where $\text{H}_2$ observed right at catalyst entrance		
Catalyst support modification	Monolithic support	Syngas formation on Pd followed indirect route where $\text{H}_2$ formed by SMR and WGS after complete $\text{O}_2$ consumption		
Catalyst support modification	Monolithic support	Co–Ni–Ru turned out as the best catalyst from the list of Co, Co–Ni, Co–Ru, Co–Ni–Ru and Ni catalysts loaded on cordierite monolith at 600°C–850°C, which demonstrated selectivity of 93.10% $\text{H}_2$ and 93.81% CO, and $\text{CH}_4$ conversion of 98.71%	General demonstration of comparison in reaction performance improvement	Figen and Baykara 2015
Catalyst support modification	Monolithic support	Pt on a FeCr alloy woven metal support effectively oxidized $\text{CH}_4$ in the presence of $\text{O}_2$ /Ar mixtures, due to its high conductivity		
Catalyst support modification	Monolithic support	Better syngas selectivity than on commercial Pt gauzes at 900°C and 1 bar and 20 bar, $\text{CH}_4$ conversion reached thermodynamic equilibrium at 20 bar	General demonstration of comparison in reaction performance improvement	Neagoe et al. 2016

Table 3 (continued)

Area of study	Investigation	Finding	Achievement	Investigator
New reactor configuration	Rh catalyst bed embedded in a Swiss-roll structured reactor, simulated using a finite-volume method with the SIMPLE algorithm by ANSYS FLUENT v12	Improved $\text{CH}_4$ conversion and $\text{H}_2$ yield by > 10% due to enhancement in combustion, steam reforming and dry reforming of $\text{CH}_4$ through waste heat recovered by reactants from product gas POX performance improved with the increase in the number of turns in the reactor or lower GHSV of reactants Variations of the $\text{H}_2\text{O}-\text{CH}_4$ ratio and carrier gas ratio with amplitude of 0.3–0.5 improved $\text{H}_2$ selectivity and $\text{CH}_4$ conversion by 1.3 and 1.1 times, respectively, compared to steady state operation	POX in a Swiss-roll reactor structure achieved high $\text{H}_2$ yield while improved energy efficiency of the system	Chen et al. 2010
Process parameters optimization	Numerical investigation on periodic feed to Pd-membrane reactor instead of steady state feed		Theoretical evidence of forced periodic operation as a beneficial condition for reaction performance improvement	Chibane et al. 2011
Reactor design modification	Substitution of a conventional dense membrane with a three-layer structure perovskite oxide membrane (a dense thin film + two porous layers deposited with Ni)	$\text{Ba}_{0.9}\text{Co}_{0.7}\text{Fe}_{0.2}\text{Nb}_{0.1}\text{O}_{3-6}$ catalytic membrane (Ni-BCFN9721) demonstrated higher $\text{H}_2$ selectivity over CO due to oxidation of dissociated carbon; higher $\text{CH}_4$ conversion due to high oxygen permeation flux; lower carbon deposition with no apparent degradation of perovskite structure	Higher performance and stability membrane in comparison to a conventional dense membrane	Song et al. 2015
Reactor design modification and process parameters optimization	FLUENT simulation and experiment of a multi-layer asymmetric oxygen-permeable membrane	Membrane based on macroporous Ni-Al foam substrate with three layers of perovskite-fluorite nanocomposites with meso-micro porosity, thin dense $\text{MnFe}_2\text{O}_4-\text{Ce}_{0.9}\text{Gd}_{0.1}\text{O}_{2-x}$ layer and catalytic porous layer of $\text{LaNi}_{0.9}\text{Pt}_{0.1}\text{O}_{3/2}/\text{Pr}_{0.3}\text{Ce}_{0.35}\text{Zr}_{0.35}\text{O}_{2-x}$ at $900^\circ\text{C}$ , varied $\text{CH}_4$ inlet concentrations of 1–20 vol. %, $\text{CH}_4$ flow rates of 1–7 l/h, air flow rates of 0.7–3.6 l/h Simulation validated experimentally with standard deviation < 12%	Optimization of process parameters (i.e. temperature, inlet gas flow rates) and membrane parameters (i.e. pore diameter in porous layer and the layer's thickness) for highest $\text{CH}_4$ conversion and syngas selectivity	Shelepova et al. 2016
Reactor design configuration	Ceramic ion transport membranes that conduct oxygen ions at elevated temperatures	Demonstration of a single compact ceramic membrane reactor that combined air separation and high-temperature syngas generation processes	Potential capital investment reduction for $\text{H}_2$ or syngas production plants	Dyer et al. 2000, Miller et al. 2014
New reactor configuration	Simulations of air injection SFR, FBR and MR	Optimized number of air injections in SFR up to 3 demonstrated an increase in $\text{H}_2$ selectivity from 50% to 53% when compared with MR Single air injection in SFR decreased the outlet temperature to 1078 K from 1269 K in FBR	Creation of a process with controllable operating temperature that eliminated the problem of high operating temperature in FBR and reduced the high operating cost in MR	Dashliborun et al. 2013
New reactor configuration	Microchannel reactor made of Ni-Pt/LaCeZrO catalyst deposited on FeCrAlloy microchannel plates	One microchannel plate demonstrated 82% $\text{CH}_4$ conversion and 60% CO selectivity at GHSV = 820,000 $\text{h}^{-1}$ and thermocycling of up to $900^\circ\text{C}$	$\text{H}_2$ production proportional to the number of microchannel plates	Makarshin et al. 2015

Table 3 (continued)

Area of study	Investigation	Finding	Achievement	Investigator
Process design modification	Modeling study of oxygen activation by electric discharge of methane-steam blend in a flow reactor	High syngas yield up to 89% by POX at the $\text{CH}_4/\text{O}_2$ ratio of 3; highest $\text{H}_2$ selectivity over CO of 4 : 1 at syngas yield of 69–76%. Steam addition delayed the conversion process and halted the increase of $\text{H}_2$ selectivity at the reactor exit	Energy released from POX is used to produce non-equilibrium plasma comprising reactive atoms/molecules for reactant preheating, hence the energy-efficient process demonstrated with high energy returned per energy invested	Starik et al. 2015
Process design modification	Combined co-electrolysis with POX	Based on the reacted $\text{CH}_4$ and the energy required for heating up $\text{CH}_4$ , efficiency of the POX-assisted co-electrolysis showed 57% improvement (from 0.21 to 0.33 at 200 mA/cm <sup>2</sup> ) as compared to conventional solid oxide $\text{H}_2\text{O}/\text{CO}_2$ co-electrolysis without POX, on top of using lower electric input	Energy-efficient promotion effect by POX on the co-electrolysis process for establishment of a clean and effective energy conversion system	Wang et al. 2015

tested operation temperatures, while the catalysts maintained high stability during the time on stream. The study demonstrated that the feed ratio played a significant role on the catalytic activity where the increase in the oxygen-methane ratio up to 0.5, and in the steam-methane ratio up to 1, improved the methane conversion.

Catalytic investigations of ATR of methane have also included the use of bimetallic catalysts, where the second metal acts as a promoter to improve the structural and catalytic properties and hence enhance methane conversion and product yield and selectivity. A study by Sepehri et al. (2016) demonstrated the effect of adding small amounts (~3 wt%) of La, Ce, Sr and Zr as a promoter on the catalytic activity of nickel catalysts supported on  $\gamma\text{-Al}_2\text{O}_3$  in ATR of methane. In general, the addition of the promoters investigated in this work resulted in high catalytic stability, without any formation of carbon on the catalyst surfaces. The addition of Zr helped improve the nickel catalyst reducibility, while the addition of Sr, Ce and La promoted greater dispersion of nickel on  $\gamma\text{-Al}_2\text{O}_3$ , with Ce exhibiting the highest activity.

Ismagilov et al. (2014a) examined the effects of metal promoter type that included Pt, Pd, Re, Mo, Sn added to Ni supported on  $\text{La}_2\text{O}_3$  at a varying promoter/Ni molar ratio of between 0.01 and 0.05 in the ATR of methane at low reaction temperatures of around 700–800°C. A different promoter would impose a different level of impact on Ni species reducibility. At a promoter/Ni molar ratio of 0.01, it was demonstrated that the order of promoters that increase methane conversion and product ( $\text{H}_2$ , CO) selectivity was arranged in the manner of  $\text{Pt} < \text{Sn} < \text{Mo} < \text{Re} < \text{Pd}$ .

On the basis of the above result, Ismagilov et al. (2014b) expanded their investigation by studying the influence of  $\text{Ce}_{0.5}\text{Zr}_{0.5}\text{O}_2$  and  $\text{La}_2\text{O}_3$  additives on the physicochemical properties of Ni-Pd catalysts supported on alumina, using different catalyst preparation techniques, incipient wetness co-impregnation and sequential impregnation. The performance difference between  $\text{Ni-Pd}/\text{Ce}_{0.5}\text{Zr}_{0.5}\text{O}_2/\text{Al}_2\text{O}_3$  and  $\text{Ni-Pd}/\text{La}_2\text{O}_3/\text{Ce}_{0.5}\text{Zr}_{0.5}\text{O}_2/\text{Al}_2\text{O}_3$  was analyzed, and the best methane ATR performance was exhibited by  $10\text{Ni}_{0.5}\text{Pd}/10\text{Ce}_{0.5}\text{Zr}_{0.5}\text{O}_2/\text{Al}_2\text{O}_3$ , with Ni-Pd catalysts prepared by the sequential impregnation technique. It was concluded that different support compositions have a different degree of Ni-support interaction, and hence contribute to the homogeneity and uniformity of NiO catalyst dispersion, and Ni species reduction ability.

Researchers have also expanded the horizon of possible performance improvement in ATR through modification in a catalyst support substrate. The performance comparison between two substrates, structured ceramic



honeycomb monolith and open cell foams, supporting Ni catalyst in methane ATR was done by Ciambelli et al. (2010). The reactor was integrated with a heat exchanger to exchange the heat from the exhaust stream to the inlet water and air streams in order to establish a process that was totally self-sustainable in the compact system. The steam-methane ratio was fixed at 0.49 and the GHSV was kept constant at  $12,300 \text{ h}^{-1}$ , while the oxygen-methane ratio was varied between 0.56 and 0.75. Foam structured catalyst exhibited higher activity (Ciambelli et al. 2010). A subsequent study (Palma et al. 2012) demonstrated equally good thermal conductivity and low average temperature with uniform temperature profile at the axial length of the catalytic bed in both monolithic support systems. Whilst this finding is promising toward commercialization of the foam-supported catalyst as a fuel cell, further investigation work at higher space velocity is still required since further analysis by Palma et al. (2012) revealed that only the first zone of the catalytic volume in the reactor bed was exploited, based on the analysis of temperature and gas composition along the bed length.

Table 4 summarizes the knowledge and information gained on ways to enhance reaction performance from the work performed by the researchers on the ATR, as reviewed in this paper.

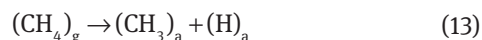
### 3.3.4 Methane decomposition

Catalytic decomposition (or cracking) of methane offers potential route to production of high purity hydrogen, due to elimination of the water-gas shift reaction in the process, hence, removing the requirement for  $\text{CO}_2$  abstraction stages. This gives a direct impact on carbon credit (Mondal and Chandran 2014). A further purification step is not required for the  $\text{CO}_x$ -free hydrogen produced by thermocatalytic decomposition, and hence can directly be used as the fuel in  $\text{H}_2$ - $\text{O}_2$  fuel cell and internal combustion engine. This laboratory-level technology requires further intensive research and development before it can be scaled up to the industrial scale (Ashik et al. 2015).

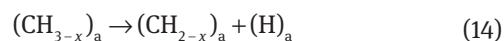
In addition, the decomposition process produces carbon nanostructure as co-product, a novel nanomaterial that offers properties such as good mechanical strength, high surface area and great thermal conductivity, desired for applications as gas storage, additives in polymer, support for catalyst or the catalyst itself, to name a few (Teo et al. 2003, Ashik et al. 2015).

The reaction mechanism involves following five steps as described by Ashik et al. (2015):

- (i) Chemisorption of methane on the leading (front-exposed) face of a catalyst particle.
- (ii) Detachment of a chemisorbed methane molecule through breaking of C–H bonds:



followed by the release of elemental carbon and hydrogen through a series of surface stepwise dissociation reactions:



where  $0 < x < 2$ ; subscripts (a) and (g) symbolize adsorbed and gaseous species, respectively.

- (iii) Adsorption of atomic hydrogen aggregation into molecules, followed by emission of gas phase:



- (iv) Aggregation of atomic carbon into encapsulated carbon by concentration difference, which leads to atomic carbon diffusion through the catalyst particle.
- (v) Nucleation of carbon and subsequent growth of carbon filament in the trailing face of the catalyst particle:



where subscript (c) denotes crystalline species.

This mechanism is derived based on the growth model postulated by Baker (1989) (Figure 6), which has become the most commonly accepted mechanism (Teo et al. 2003) for explaining CNF growth by catalytic decomposition of hydrocarbon. The model was proposed based on the synthesis of CNFs for various potential applications, which is a separate subject of interest.

Ideas in terms of process design have been evaluated to address the challenge in separating the carbon nanostructure deposited on the catalyst during the decomposition process, from the catalyst surface, in a continuous mode. This challenging requirement to separate carbon from the catalyst surface has the potential to be overcome based on the findings in a simulated study conducted by Mondal and Chandran (2014). The investigation explored low-temperature hydrogen production from a stepwise cyclic process of thermo-catalytic decomposition of methane ( $\text{CH}_4 \rightarrow 2\text{H}_2 + \text{C}$ ), followed by gasification by steam ( $\text{C} + 2\text{H}_2\text{O} \rightarrow 2\text{H}_2 + \text{CO}_2$ ) at the same temperature. In addition, this stepwise cyclic process produced hydrogen slightly higher than that from the conventional steam reforming process for every mole of methane, on top of operating at much lower temperature ( $600^\circ\text{C}$ ), hence saving significant energy. In the cyclic process, the re-oxidation of the metal catalysts took place in the

Table 4: Investigation areas in methane autothermal reaction to improve reaction performance.

Area of investigation	Investigation	Finding	Achievement	Investigator
Process parameters optimization	Manipulation of operation parameters	MATLAB simulation: SMR prone to backward at low temperature; there was an inflection point temperature that the reaction moved forward At $H_2O-CH_4 = 2$ : Air- $CH_4 = 1$ kept the inflection point temperature at 700–800 K; an increase in air- $CH_4$ brought up the inflection point temperature At > 1000 K: increase in air- $CH_4$ decreased $H_2$ yield Optimized parameters for ATR demonstrated at 1000 K, air- $CH_4 = 1$ and $H_2O-CH_4 = 2$ Regulation of $H_2O-CH_4$ manipulated selectivity of $H_2/CO$	Knowledge on techniques to improve $H_2/CO$ selectivity and increase $CH_4$ conversion	Yan et al. 2013
Process parameters optimization	Manipulation of operation parameters	CHEMKIN simulation of ATR in a micro-reactor for mobile fuel cell development Optimum conditions at $O_2-CH_4 = 0.4$ , $H_2O-CH_4 = 2.1$ , temperature = 1073 K, flow velocity = 1 m/s, catalyst active site density = $9.98 \times 10^{-6}$ kmol/m <sup>2</sup> , achieved $CH_4$ conversion = 93–99%, $H_2$ yield = 65–80%, CO yield = 50–65%, carbon deposition = $1.65 \times 10^{-12}$ kmol/m <sup>2</sup> Highest $CH_4$ conversion observed on 15% Ni/MgAl <sub>2</sub> O <sub>4</sub> at temperatures = 550–700°C $O_2-CH_4$ ratio up to 0.5 and $H_2O-CH_4$ up to 1 improved $CH_4$ conversion further Addition of Zr improved Ni reducibility, addition of Sr, Ce and La promoted greater Ni dispersion on $\gamma-Al_2O_3$ , Ce exhibited the highest activity at operating temperatures 550–700°C Effects of promoter type and molar ratio Me/Ni (Me = Pt, Pd, Re, Mo, Sn; Me/Ni = 0.01–0.05) on Ni-Me/La <sub>2</sub> O <sub>3</sub> in terms of structural, reducing and catalytic properties At 700–800°C, Me/Ni = 0.01: $CH_4$ conversion and syngas yield increased in order of Pt < Sn < Mo < Re < Pd, correlated with increasing reducibility of Ni species At 850°C, LaNi <sub>0.99</sub> Pd <sub>0.01</sub> O <sub>3</sub> catalyst demonstrated ~100% $CH_4$ conversion, ~41% $H_2$ and ~57% CO	Knowledge on techniques to improve $H_2/CO$ selectivity and increase $CH_4$ conversion at minimal carbon deposition	Yan et al. 2015
Catalyst loading amount optimization	Varied nickel content (5, 10 and 15%) supported on MgAl <sub>2</sub> O <sub>4</sub>		Information on optimal Ni content and feed ratios for the best catalytic activity	Mosayebi et al. 2012
Catalyst modification	Effect of adding small amounts (~3 wt%) of La, Ce, Sr and Zr as promoter on catalytic activity of Ni/ $\gamma-Al_2O_3$		Information on the rank of promoters in improving catalytic activity	Sepehri et al. 2016
Catalyst modification	Development of bimetallic catalysts Ni-Me/La <sub>2</sub> O <sub>3</sub> (Me = Pt, Pd, Re, Mo, Sn)		Successful selection of the second active metal added to Ni to enhance catalytic activity in ATR	Ismagilov et al. (2014a)

Table 4 (continued)

Area of investigation	Investigation	Finding	Achievement	Investigator
Catalyst support modification and preparation technique	Influence of additives ( $\text{Ce}_{0.5}\text{Zr}_{0.5}\text{O}_2$ versus $\text{La}_{2.3}/\text{Ce}_{0.5}\text{Zr}_{0.5}\text{O}_2$ ), and influence of catalyst preparation techniques (incipient wetness co-impregnation versus sequential impregnation) on physicochemical properties of Ni-Pd/ $\text{Al}_2\text{O}_3$	Addition of additive provided high $\text{CH}_4$ conversion (90–100%), CO yield (55–85%) and $\text{H}_2$ yield (55–75%) at 750–950°C Best catalytic performance provided by 10 Ni <sub>0.5</sub> Pd/10Ce <sub>0.5</sub> Zr <sub>0.5</sub> O <sub>2</sub> /Al <sub>2</sub> O <sub>3</sub> catalyst prepared by the Pd-Ni sequential impregnation technique that resulted in peculiar Ni-Pd particles structure At $\text{H}_2\text{O}/\text{CH}_4 = 0.49$ , GHSV = 12,300 h <sup>-1</sup> , O <sub>2</sub> /CH <sub>4</sub> = 0.56–0.75, foam-structured catalysts exhibited higher activity and lower average temperature along the axial length with more uniform profile	Preparation method determines the support composition and catalyst reducibility Support composition influences the Ni-support interaction strength that promotes NiO dispersion and reducibility of Ni species	Ismagilov et al. 2014b
Catalyst support modification	Structured ceramic honeycomb monolith versus open cell foams, supporting Ni catalyst in reactor integrated with heat exchanger to exchange heat from exhaust stream to inlet water and air for the self-sustainable process		Development of a compact reactor with a thermally self-sustainable system for fuel cell application	Ciambelli et al. 2010, Palma et al. 2012

regeneration step and then the particle was re-reduced in  $\text{H}_2$  for reactivation, and the methane decomposition step repeated accordingly.

Methane is one of the most inactive hydrocarbons that has very strong carbon-hydrogen bond and high symmetrical molecular structure. In the absence of a catalyst, its efficient decomposition could only occur at temperatures beyond 1200°C in order to break the strong carbon-hydrogen bond (440 kJ/mol) due to its high thermodynamic stability. There have been various studies conducted in the attempt to bring down this temperature through utilization of metal and carbon-derived catalysts (Ashik et al. 2015). A large amount of amorphous carbon is formed during this endothermic reaction that leads to catalyst deactivation. Various studies have been carried out to overcome this issue, through modifications of transitional metals, rare earth (RE) metals, alloys and carbon material catalysts.

Bayat et al. (2016b) presented a catalytic performance study on thermocatalytic decomposition of methane using varied nickel catalyst loading amounts on mesoporous nanocrystalline  $\gamma\text{-Al}_2\text{O}_3$  support. The level of catalytic activity changed with the variation in nickel content, whereas the stability of the catalysts was dependent on the reaction temperature. Nickel content at 50 wt% demonstrated the highest catalyst's activity and stability at reaction temperatures of 625°C and 650°C, but the reaction rates increased and accelerated the catalyst deactivation when the reaction was run at higher operating temperatures. It was also noted that an operation at lower temperatures decreased the methane conversion. The morphology of the used catalysts from scanning electron microscopy analysis revealed formation of CNFs with diameter close to the nickel particle size, but the diameter of the CNFs decreased with the increase in the operating temperature.

Transitional metals other than nickel have also been used in the methane decomposition performance studies. Ibrahim et al. (2015) investigated the activity performance by different iron loadings of 15–100% supported on alumina catalyst. It was revealed that hydrogen yield increased with higher iron loading up to 60% where loadings higher than this were found to have decreased the catalyst's surface area. At 700°C, the yield of hydrogen over 60% Fe/ $\text{Al}_2\text{O}_3$  catalyst was 77%.

Investigation on the addition of a second transitional metal to the first has also been carried out. Bayat et al. (2016c) examined the catalytic behavior of iron added to nickel catalysts on  $\gamma$ -alumina support. Analysis conducted confirmed formation of  $\text{NiFe}_2\text{O}_4$  species in calcined Ni-Fe alloy catalysts, but the degree of reducibility reduced with increasing composition of iron in the alloy. However, the addition of iron improved the catalytic stability through

enhancement of the carbon diffusion rate (step C in Figure 6) and prevention of carbon encapsulation that deactivates the catalyst at reaction temperatures of below 650°C.

Bayat et al. (2016a) expanded the study by carrying out a catalytic performance investigation in methane decomposition using 50 wt% Ni supported on alumina, with an addition of fixed Fe content of 10% and varied Cu contents of 0, 5, 10 and 15%, as promoters. Although it was found that iron decreased the reducibility of the nickel catalyst, copper on the other hand compensated this effect by increasing the adsorption of methane while at the same time improving the dispersion of nickel particles on the support. It was revealed that the addition of both promoters resulted in an improvement in the catalytic performance, due to an enhancement of carbon diffusion rate that helped prevent coke formation on the nickel surface. Additionally, the high affinity of copper with a graphite structure contributed to further effect of coke formation prevention, which slowed down the catalyst deactivation at reaction temperatures of above 700°C, with 50%Ni-10%Fe-10%Cu/Al<sub>2</sub>O<sub>3</sub> displaying optimum catalytic performance.

In an earlier work, the catalytic behavior of Ni-Cu-containing catalyst particles was investigated to understand the effect of adding additional transitional metal. Different compositions of Ni-Cu-Co alloy catalysts prepared by Lua and Wang (2014) were observed to have resulted in porous aggregates of Ni-Cu-Co alloy particles of different

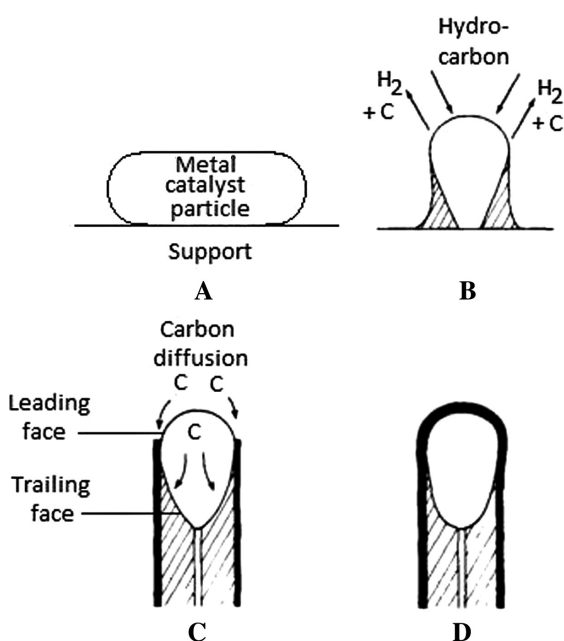
crystalline sizes. Based on general observation on the tri-metal alloy, good catalytic activities on methane decomposition were exhibited at temperatures between 650 and 775°C. Due to higher melting point of cobalt, the addition of it resulted in effective inhibition of a quasi-liquid phenomenon at higher temperatures, and hence, improved catalysts stability, in comparison with the Ni-Cu bi-metallic alloy. However, a continued increase in cobalt content led to phase separation.

A study on the effect of cobalt content was also carried out by Awadallah and Aboul-Enein (2015) over bimetallic catalysts containing a series of Co/W at 40/10, 30/20, 20/30 and 10/40 wt%, supported on MgO with a total metal content of 50 wt%. It was revealed that the 40%Co-10%W/MgO catalyst exhibited the highest activity for the production of hydrogen as well as the carbon nanostructure (observed in the form of multi-walled carbon nanotubes in this study). Based on their analysis, this catalyst superiority achievement was attributed to the large amount formation of non-interacted Co<sub>3</sub>O<sub>4</sub> species.

In general, a lot of other investigations in methane decomposition catalytic activity are focusing on the effect of adding the content of other metals on nickel-based catalyst. Ying et al. (2014) synthesized different RE-Ni bimetallic catalysts to evaluate the performance of methane decomposition in a fixed bed microreactor. The RE utilized in their study included La, Ce, Pr, Nd, Sm and Y, where it was demonstrated that the RE-modified nickel catalysts resulted in smaller nickel particle size, good thermal stability and high activity, on top of preventing solid carbon formation. Of all the REs, La was found to be the best additive in the examination.

Carbon materials have also been seen as promising catalysts for methane decomposition and hence serve as an alternative to expensive transition metals. Activated carbon (AC) is able to provide reasonable reaction kinetics for production of continuous high purity hydrogen, as investigated by Liu et al. (2016) through the application of a chemical looping process. No signs of deactivation and agglomeration of AC catalyst were observed during the cyclic study under proper regeneration, and CO<sub>2</sub>-free hydrogen of high purity was produced continuously. The application of AC catalyst is enabled by its crushing strength, and hence, it is suitable for a chemical looping process.

Shen and Lua (2016) presented methods to synthesize porous carbon of an interconnected trimodal pore system as the catalyst for decomposition of methane. The trimodal pore system that was synthesized consisted of intra-mesopores and micropores that provided a large surface area for reactant accessibility to active sites and



**Figure 6:** Growth mechanism of carbon nanofibers by catalytic hydrocarbon decomposition postulated by Baker (1989).



Table 5: Investigation areas in methane decomposition and their corresponding benefits.

Area of investigation	Investigation	Finding	Achievement	Investigator
Process design	Aspen Plus simulation of a stepwise cyclic process of methane decomposition and steam gasification in series versus conventional SMR	At 600°C (significantly lower than SMR operating temperature), re-oxidation of metal catalysts took place in the regeneration step and then the particle re-reduced in H <sub>2</sub> for reactivation, and methane decomposition step repeated	Low-temperature operation (energy saving), continuous removal of carbon deposit	Mondal and Chandran, 2014
Catalyst loading amount optimization	Varied Ni catalyst loading amounts (30, 40, 50 and 60%) on $\gamma$ -Al <sub>2</sub> O <sub>3</sub> support	Ni at 50 wt% demonstrated highest catalytic activity and stability at 625°C and 650°C At higher temperatures: reaction rates increased, hence, accelerated catalyst deactivation At lower temperatures: CH <sub>4</sub> conversion decreased	Information on optimal Ni content and operating temperatures for highest catalytic activity and stability	Bayat et al. 2016b
Catalyst loading amount optimization	Varied Fe catalyst loading amounts (15–100%) on Al <sub>2</sub> O <sub>3</sub> support	Fe at 60 wt% demonstrated maximum hydrogen yield of 77%, with further loading amounts led to a decrease in the catalyst's surface area Addition of Fe improved catalytic stability through enhancement of carbon diffusion rate and prevention of coke formation at below 650°C	Information on optimal Fe content for highest catalytic activity and hydrogen selectivity	Ibrahim et al. 2015
Catalyst modification	Influence of Fe content (0, 5, 10 and 15%) on Ni catalysts (50%) supported on $\gamma$ -Al <sub>2</sub> O <sub>3</sub>	50%Ni-10%Fe/Al <sub>2</sub> O <sub>3</sub> exhibited highest activity but an increase in reaction temperature > 700°C decreased CH <sub>4</sub> conversion due to coke formation Addition of both promoters (Fe and Cu) resulted in an improvement in the catalytic performance, due to enhancement of carbon diffusion rate that helped prevent coke formation on the Ni surface	Information on optimal Fe content in Ni catalysts for catalyst's stability enhancement	Bayat et al. 2016c
Catalyst modification	Influence of Cu content (0, 5, 10 and 15%) on Ni-Fe catalysts (Fe fixed at 10%)	High affinity of Cu with graphite structure prevented coke formation, hence, slowed down catalyst deactivation at temperatures > 700°C 50%Ni-10%Fe-10%Cu/Al <sub>2</sub> O <sub>3</sub> displayed most optimum catalytic performance	Information on optimal Cu content in Ni-Fe catalysts activity enhancement and coke formation prevention at high operating temperatures	Bayat et al. 2016a
Catalyst modification	Influence of Co content on Ni-Cu catalysts	Addition of Co led to smaller crystalline size alloy particles compared to Ni-Cu or pure Ni catalysts Increasing Co improved catalyst stability at higher temperatures (~750°C) due to higher Co melting point than Ni and Cu, but further continual Co addition led to phase separation Co-Cu could not remain stable in the reaction without Ni, and Co particles deactivated easily due to encapsulation by carbon	Information on optimal Co content in Ni-Cu catalysts for activity enhancement	Lua and Wang 2014

Table 5 (continued)

Area of investigation	Investigation	Finding	Achievement	Investigator
Catalyst modification	Influence of Co content on W catalysts at Co/W = 40/10, 30/20, 20/30 and 10/40 wt%, supported on MgO	At 700°C, 40%Co-10%W/MgO catalyst exhibited the highest activity with H <sub>2</sub> yield of 88.5%, in comparison to 79.8% by 50%Co/MgO, attributed to the large amount formation of non-interacted Co <sub>3</sub> O <sub>4</sub> species 20%Co-30%W and 10%Co-40%W demonstrated formation of almost non-reducible CoWO <sub>4</sub> and MgWO <sub>4</sub> particles, which reduced the catalytic activity	Information on optimal Co content in W catalysts for activity enhancement	Awadallah and Aboul-Enein 2015
Catalyst modification	Influence of rare earth (RE)-Ni bimetallic catalysts, RE = La, Ce, Pr, Nd, Sm and Y	RE-modified Ni catalysts resulted in smaller nickel particle size, good thermal stability, high activity, minimal, or no carbon formation La found to be the best additive in the examination	Information on the best RE as an additive to promote Ni catalysts for activity enhancement	Ying et al. 2014
Carbon material catalyst	Use of activated carbon (AC) as active cyclic material in the chemical looping process	AC exhibited good durability (good crushing strength with no agglomeration) and able to be regenerated, where active sites restored and reappeared on the surface again	Chemical looping using AC demonstrated to be feasible for continuous H <sub>2</sub> production of high purity	Liu et al. 2016
Carbon material catalyst	Use of porous carbon of an interconnected trimodal pore system	Synthesis of porous carbon that had a large surface area for reactant accessibility to active sites and spaces for deposition of carbon from the decomposition due to its intra-mesopores and micropores; and effective bulk flow of reactant molecules due to its inter-mesopores	Demonstration of catalytic activity of the synthesized trimodal porous carbon that provided higher activity than commercial carbon materials	Shen and Lua 2016

spaces for deposition of carbon from the decomposition, and inter-mesopores for effective bulk flow of the reactant molecules, and hence, minimized reactant's diffusion resistance. Higher methane conversion and hydrogen yield were achieved with as-synthesized carbon in this study compared to the commercial carbon materials.

Table 5 provides summarized findings on efforts taken in improving methane decomposition in terms of conversion efficiency, carbon deposition minimization, hydrogen purification as well as energy saving through low-temperature operation as reviewed in this paper.

## 4 Conclusions

Hydrogen finds applications in a large production scale at chemical plants and oil refineries, as a raw material for the synthesis of mainly ammonia and methanol, and as reactant in hydrocracking and hydrofining of oil to produce transportation fuels and remove contaminants such as sulfur to meet the emission legislation and fuel specifications. At a smaller scale where hydrogen is being generated from renewable energy sources like wind and solar energy that power the water electrolysis process, hydrogen finds its applications as an energy carrier being stored on electricity grids, and as fuel in hydrogen vehicles, used in internal combustion engines and fuel cells, and hence, reduces the reliance on fossil fuels and the emissions of greenhouse gas.

The main routes of hydrogen production at a large scale for industrial utilization are widely coming from thermocatalytic processing of methane, the main component in natural gas, which accounts for close to half of the total hydrogen production resource. The same technologies are also applicable to process petroleum oil components such as naphtha, which accounts for slightly less than a third of the total resource. Whilst methane decomposition or cracking technology is still at the laboratory scale level and has yet to find its place on an industrial scale, SMR dominates the hydrogen production technology, along with two other most commonly used catalytic hydrocarbon processing technologies: POX and ATR. These three technologies are the most economical and efficient for large-scale hydrogen production.

In all the technologies reviewed, a lot of interest has been developed among researchers to introduce initiatives in improving methane conversion, hydrogen selectivity and energy efficiency as well as suppressing coke formation that accelerates catalyst deactivation. Among the general efforts identified from the review include studies on qualifying various oxygen-transfer materials

as the catalyst support; addition of the second and third metals to the primary metal catalyst, or addition of mixed perovskite-like oxides to the primary support material to act as promoters; the use of monolithic catalytic support to replace the pelletized catalytic system; reactor configuration modifications; and engineering process modifications.

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